
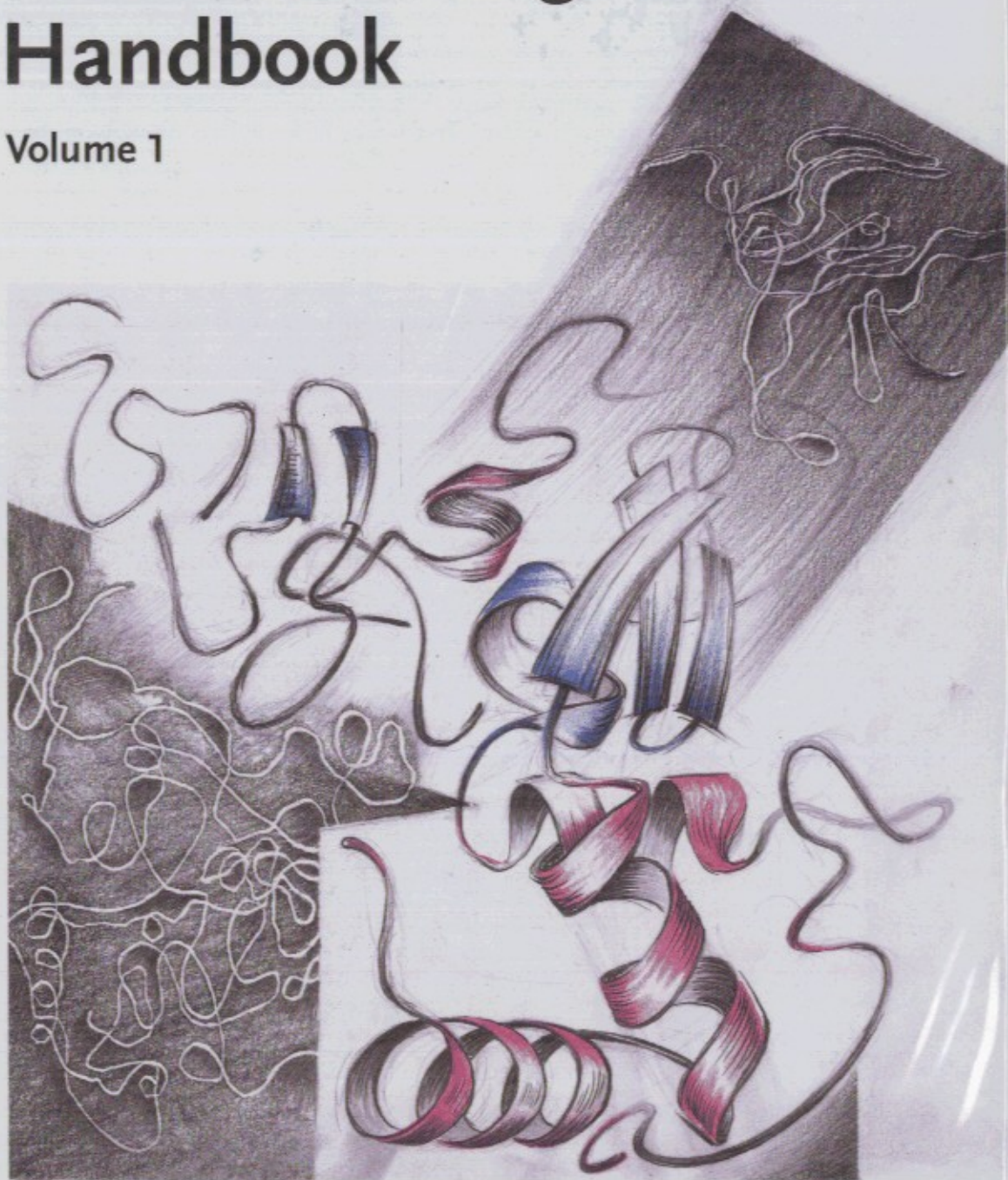


Edited by
Johannes Buchner, Thomas Kiefhaber

 WILEY-VCH

Protein Folding Handbook

Volume 1



Contents

Part I, Volume 1

Preface LVIII

Contributors of Part I LX

I/1	Principles of Protein Stability and Design	1
1	Early Days of Studying the Mechanism of Protein Folding	3
	<i>Robert L. Baldwin</i>	
1.1	Introduction	3
1.2	Two-state Folding	4
1.3	Levinthal's Paradox	5
1.4	The Domain as a Unit of Folding	6
1.5	Detection of Folding Intermediates and Initial Work on the Kinetic Mechanism of Folding	7
1.6	Two Unfolded Forms of RNase A and Explanation by Proline Isomerization	9
1.7	Covalent Intermediates in the Coupled Processes of Disulfide Bond Formation and Folding	11
1.8	Early Stages of Folding Detected by Antibodies and by Hydrogen Exchange	12
1.9	Molten Globule Folding Intermediates	14
1.10	Structures of Peptide Models for Folding Intermediates	15
	Acknowledgments	16
	References	16
2	Spectroscopic Techniques to Study Protein Folding and Stability	22
	<i>Franz Schmid</i>	
2.1	Introduction	22
2.2	Absorbance	23
2.2.1	Absorbance of Proteins	23
2.2.2	Practical Considerations for the Measurement of Protein Absorbance	27

2.2.3	Data Interpretation	29
2.3	Fluorescence	29
2.3.1	The Fluorescence of Proteins	30
2.3.2	Energy Transfer and Fluorescence Quenching in a Protein: Barnase	31
2.3.3	Protein Unfolding Monitored by Fluorescence	33
2.3.4	Environmental Effects on Tyrosine and Tryptophan Emission	36
2.3.5	Practical Considerations	37
2.4	Circular Dichroism	38
2.4.1	CD Spectra of Native and Unfolded Proteins	38
2.4.2	Measurement of Circular Dichroism	41
2.4.3	Evaluation of CD Data	42
	References	43
3	Denaturation of Proteins by Urea and Guanidine Hydrochloride	45
	<i>C. Nick Pace, Gerald R. Grimsley, and J. Martin Scholtz</i>	
3.1	Historical Perspective	45
3.2	How Urea Denatures Proteins	45
3.3	Linear Extrapolation Method	48
3.4	$\Delta G(\text{H}_2\text{O})$	50
3.5	m -Values	55
3.6	Concluding Remarks	58
3.7	Experimental Protocols	59
3.7.1	How to Choose the Best Denaturant for your Study	59
3.7.2	How to Prepare Denaturant Solutions	59
3.7.3	How to Determine Solvent Denaturation Curves	60
3.7.3.1	Determining a Urea or GdmCl Denaturation Curve	62
3.7.3.2	How to Analyze Urea or GdmCl Denaturant Curves	63
3.7.4	Determining Differences in Stability	64
	Acknowledgments	65
	References	65
4	Thermal Unfolding of Proteins Studied by Calorimetry	70
	<i>George I. Makhatadze</i>	
4.1	Introduction	70
4.2	Two-state Unfolding	71
4.3	Cold Denaturation	76
4.4	Mechanisms of Thermostabilization	77
4.5	Thermodynamic Dissection of Forces Contributing to Protein Stability	79
4.5.1	Heat Capacity Changes, ΔC_p	81
4.5.2	Enthalpy of Unfolding, ΔH	81
4.5.3	Entropy of Unfolding, ΔS	83
4.6	Multistate Transitions	84
4.6.1	Two-state Dimeric Model	85

4.6.2	Two-state Multimeric Model	86
4.6.3	Three-state Dimeric Model	86
4.6.4	Two-state Model with Ligand Binding	88
4.6.5	Four-state (Two-domain Protein) Model	90
4.7	Experimental Protocols	92
4.7.1	How to Prepare for DSC Experiments	92
4.7.2	How to Choose Appropriate Conditions	94
4.7.3	Critical Factors in Running DSC Experiments	94
	References	95
5	Pressure–Temperature Phase Diagrams of Proteins	99
	<i>Wolfgang Doster and Josef Friedrich</i>	
5.1	Introduction	99
5.2	Basic Aspects of Phase Diagrams of Proteins and Early Experiments	100
5.3	Thermodynamics of Pressure–Temperature Phase Diagrams	103
5.4	Measuring Phase Stability Boundaries with Optical Techniques	110
5.4.1	Fluorescence Experiments with Cytochrome <i>c</i>	110
5.4.2	Results	112
5.5	What Do We Learn from the Stability Diagram?	116
5.5.1	Thermodynamics	116
5.5.2	Determination of the Equilibrium Constant of Denaturation	117
5.5.3	Microscopic Aspects	120
5.5.4	Structural Features of the Pressure-denatured State	122
5.6	Conclusions and Outlook	123
	Acknowledgment	124
	References	124
6	Weak Interactions in Protein Folding: Hydrophobic Free Energy, van der Waals Interactions, Peptide Hydrogen Bonds, and Peptide Solvation	127
	<i>Robert L. Baldwin</i>	
6.1	Introduction	127
6.2	Hydrophobic Free Energy, Burial of Nonpolar Surface and van der Waals Interactions	128
6.2.1	History	128
6.2.2	Liquid–Liquid Transfer Model	128
6.2.3	Relation between Hydrophobic Free Energy and Molecular Surface Area	130
6.2.4	Quasi-experimental Estimates of the Work of Making a Cavity in Water or in Liquid Alkane	131
6.2.5	Molecular Dynamics Simulations of the Work of Making Cavities in Water	133
6.2.6	Dependence of Transfer Free Energy on the Volume of the Solute	134
6.2.7	Molecular Nature of Hydrophobic Free Energy	136

6.2.8	Simulation of Hydrophobic Clusters	137
6.2.9	ΔC_p and the Temperature-dependent Thermodynamics of Hydrophobic Free Energy	137
6.2.10	Modeling Formation of the Hydrophobic Core from Solvation Free Energy and van der Waals Interactions between Nonpolar Residues	142
6.2.11	Evidence Supporting a Role for van der Waals Interactions in Forming the Hydrophobic Core	144
6.3	Peptide Solvation and the Peptide Hydrogen Bond	145
6.3.1	History	145
6.3.2	Solvation Free Energies of Amides	147
6.3.3	Test of the Hydrogen-Bond Inventory	149
6.3.4	The Born Equation	150
6.3.5	Prediction of Solvation Free Energies of Polar Molecules by an Electrostatic Algorithm	150
6.3.6	Prediction of the Solvation Free Energies of Peptide Groups in Different Backbone Conformations	151
6.3.7	Predicted Desolvation Penalty for Burial of a Peptide H-bond	153
6.3.8	Gas–Liquid Transfer Model	154
	Acknowledgments	156
	References	156
7	Electrostatics of Proteins: Principles, Models and Applications	163
	<i>Sonja Braun-Sand and Arieh Warshel</i>	
7.1	Introduction	163
7.2	Historical Perspectives	163
7.3	Electrostatic Models: From Microscopic to Macroscopic Models	166
7.3.1	All-Atom Models	166
7.3.2	Dipolar Lattice Models and the PDL D Approach	168
7.3.3	The PDL D/S-LRA Model	170
7.3.4	Continuum (Poisson-Boltzmann) and Related Approaches	171
7.3.5	Effective Dielectric Constant for Charge–Charge Interactions and the GB Model	172
7.4	The Meaning and Use of the Protein Dielectric Constant	173
7.5	Validation Studies	176
7.6	Systems Studied	178
7.6.1	Solvation Energies of Small Molecules	178
7.6.2	Calculation of pK_a Values of Ionizable Residues	179
7.6.3	Redox and Electron Transport Processes	180
7.6.4	Ligand Binding	181
7.6.5	Enzyme Catalysis	182
7.6.6	Ion Pairs	183
7.6.7	Protein–Protein Interactions	184
7.6.8	Ion Channels	185
7.6.9	Helix Macrodipoles versus Localized Molecular Dipoles	185
7.6.10	Folding and Stability	186
7.7	Concluding Remarks	189

Acknowledgments	190
References	190

8 Protein Conformational Transitions as Seen from the Solvent: Magnetic Relaxation Dispersion Studies of Water, Co-solvent, and Denaturant Interactions with Nonnative Proteins 201

Bertil Halle, Vladimir P. Denisov, Kristofer Modig, and Monika Davidovic

8.1	The Role of the Solvent in Protein Folding and Stability	201
8.2	Information Content of Magnetic Relaxation Dispersion	202
8.3	Thermal Perturbations	205
8.3.1	Heat Denaturation	205
8.3.2	Cold Denaturation	209
8.4	Electrostatic Perturbations	213
8.5	Solvent Perturbations	218
8.5.1	Denaturation Induced by Urea	219
8.5.2	Denaturation Induced by Guanidinium Chloride	225
8.5.3	Conformational Transitions Induced by Co-solvents	228
8.6	Outlook	233
8.7	Experimental Protocols and Data Analysis	233
8.7.1	Experimental Methodology	233
8.7.1.1	Multiple-field MRD	234
8.7.1.2	Field-cycling MRD	234
8.7.1.3	Choice of Nuclear Isotope	235
8.7.2	Data Analysis	236
8.7.2.1	Exchange Averaging	236
8.7.2.2	Spectral Density Function	237
8.7.2.3	Residence Time	239
8.7.2.4	^{19}F Relaxation	240
8.7.2.5	Coexisting Protein Species	241
8.7.2.6	Preferential Solvation	241
	References	242

9 Stability and Design of α -Helices 247

Andrew J. Doig, Neil Errington, and Teuku M. Iqbalsyah

9.1	Introduction	247
9.2	Structure of the α -Helix	247
9.2.1	Capping Motifs	248
9.2.2	Metal Binding	250
9.2.3	The 3_{10} -Helix	251
9.2.4	The π -Helix	251
9.3	Design of Peptide Helices	252
9.3.1	Host–Guest Studies	253
9.3.2	Helix Lengths	253
9.3.3	The Helix Dipole	253
9.3.4	Acetylation and Amidation	254
9.3.5	Side Chain Spacings	255

9.3.6	Solubility	256
9.3.7	Concentration Determination	257
9.3.8	Design of Peptides to Measure Helix Parameters	257
9.3.9	Helix Templates	259
9.3.10	Design of 3_{10} -Helices	259
9.3.11	Design of π -helices	261
9.4	Helix Coil Theory	261
9.4.1	Zimm-Bragg Model	261
9.4.2	Lifson-Roig Model	262
9.4.3	The Unfolded State and Polyproline II Helix	265
9.4.4	Single Sequence Approximation	265
9.4.5	N- and C-Caps	266
9.4.6	Capping Boxes	266
9.4.7	Side-chain Interactions	266
9.4.8	N1, N2, and N3 Preferences	267
9.4.9	Helix Dipole	267
9.4.10	3_{10} - and π -Helices	268
9.4.11	AGADIR	268
9.4.12	Lomize-Mosberg Model	269
9.4.13	Extension of the Zimm-Bragg Model	270
9.4.14	Availability of Helix/Coil Programs	270
9.5	Forces Affecting α -Helix Stability	270
9.5.1	Helix Interior	270
9.5.2	Caps	273
9.5.3	Phosphorylation	276
9.5.4	Noncovalent Side-chain Interactions	276
9.5.5	Covalent Side-chain interactions	277
9.5.6	Capping Motifs	277
9.5.7	Ionic Strength	279
9.5.8	Temperature	279
9.5.9	Trifluoroethanol	279
9.5.10	pK_a Values	280
9.5.11	Relevance to Proteins	281
9.6	Experimental Protocols and Strategies	281
9.6.1	Solid Phase Peptide Synthesis (SPPS) Based on the Fmoc Strategy	281
9.6.1.1	Equipment and Reagents	281
9.6.1.2	Fmoc Deprotection and Coupling	283
9.6.1.3	Kaiser Test	284
9.6.1.4	Acetylation and Cleavage	285
9.6.1.5	Peptide Precipitation	286
9.6.2	Peptide Purification	286
9.6.2.1	Equipment and Reagents	286
9.6.2.2	Method	286
9.6.3	Circular Dichroism	287
9.6.4	Acquisition of Spectra	288

9.6.4.1	Instrumental Considerations	288
9.6.5	Data Manipulation and Analysis	289
9.6.5.1	Protocol for CD Measurement of Helix Content	291
9.6.6	Aggregation Test for Helical Peptides	291
9.6.6.1	Equipment and Reagents	291
9.6.6.2	Method	292
9.6.7	Vibrational Circular Dichroism	292
9.6.8	NMR Spectroscopy	292
9.6.8.1	Nuclear Overhauser Effect	293
9.6.8.2	Amide Proton Exchange Rates	294
9.6.8.3	^{13}C NMR	294
9.6.9	Fourier Transform Infrared Spectroscopy	295
9.6.9.1	Secondary Structure	295
9.6.10	Raman Spectroscopy and Raman Optical Activity	296
9.6.11	pH Titrations	298
9.6.11.1	Equipment and Reagents	298
9.6.11.2	Method	298
	Acknowledgments	299
	References	299
10	Design and Stability of Peptide β-Sheets	314
	<i>Mark S. Searle</i>	
10.1	Introduction	314
10.2	β -Hairpins Derived from Native Protein Sequences	315
10.3	Role of β -Turns in Nucleating β -Hairpin Folding	316
10.4	Intrinsic ϕ, ψ Propensities of Amino Acids	319
10.5	Side-chain Interactions and β -Hairpin Stability	321
10.5.1	Aromatic Clusters Stabilize β -Hairpins	322
10.5.2	Salt Bridges Enhance Hairpin Stability	325
10.6	Cooperative Interactions in β -Sheet Peptides: Kinetic Barriers to Folding	330
10.7	Quantitative Analysis of Peptide Folding	331
10.8	Thermodynamics of β -Hairpin Folding	332
10.9	Multistranded Antiparallel β -Sheet Peptides	334
10.10	Concluding Remarks: Weak Interactions and Stabilization of Peptide β -Sheets	339
	References	340
11	Predicting Free Energy Changes of Mutations in Proteins	343
	<i>Raphael Guerois, Joaquim Mendes, and Luis Serrano</i>	
11.1	Physical Forces that Determine Protein Conformational Stability	343
11.1.1	Protein Conformational Stability [1]	343
11.1.2	Structures of the N and D States [2–6]	344
11.1.3	Studies Aimed at Understanding the Physical Forces that Determine Protein Conformational Stability [1, 2, 8, 19–26]	346
11.1.4	Forces Determining Conformational Stability [1, 2, 8, 19–27]	346

11.1.5	Intramolecular Interactions	347
11.1.5.1	van der Waals Interactions	347
11.1.5.2	Electrostatic Interactions	347
11.1.5.3	Conformational Strain	349
11.1.6	Solvation	350
11.1.7	Intramolecular Interactions and Solvation Taken Together	350
11.1.8	Entropy	351
11.1.9	Cavity Formation	352
11.1.10	Summary	353
11.2	Methods for the Prediction of the Effect of Point Mutations on in vitro Protein Stability	353
11.2.1	General Considerations on Protein Plasticity upon Mutation	353
11.2.2	Predictive Strategies	355
11.2.3	Methods	356
11.2.3.1	From Sequence and Multiple Sequence Alignment Analysis	356
11.2.3.2	Statistical Analysis of the Structure Databases	356
11.2.3.3	Helix/Coil Transition Model	357
11.2.3.4	Physicochemical Method Based on Protein Engineering Experiments	359
11.2.3.5	Methods Based only on the Basic Principles of Physics and Thermodynamics	364
11.3	Mutation Effects on in vivo Stability	366
11.3.1	The N-terminal Rule	366
11.3.2	The C-terminal Rule	367
11.3.3	PEST Signals	368
11.4	Mutation Effects on Aggregation	368
	References	369

I/2 Dynamics and Mechanisms of Protein Folding Reactions 377

12.1	Kinetic Mechanisms in Protein Folding	379
	<i>Annett Bachmann and Thomas Kiefhaber</i>	
12.1.1	Introduction	379
12.1.2	Analysis of Protein Folding Reactions using Simple Kinetic Models	379
12.1.2.1	General Treatment of Kinetic Data	380
12.1.2.2	Two-state Protein Folding	380
12.1.2.3	Complex Folding Kinetics	384
12.1.2.3.1	Heterogeneity in the Unfolded State	384
12.1.2.3.2	Folding through Intermediates	388
12.1.2.3.3	Rapid Pre-equilibria	391
12.1.2.3.4	Folding through an On-pathway High-energy Intermediate	393
12.1.3	A Case Study: the Mechanism of Lysozyme Folding	394
12.1.3.1	Lysozyme Folding at pH 5.2 and Low Salt Concentrations	394
12.1.3.2	Lysozyme Folding at pH 9.2 or at High Salt Concentrations	398
12.1.4	Non-exponential Kinetics	401

12.1.5	Conclusions and Outlook	401
12.1.6	Protocols – Analytical Solutions of Three-state Protein Folding Models	402
12.1.6.1	Triangular Mechanism	402
12.1.6.2	On-pathway Intermediate	403
12.1.6.3	Off-pathway Mechanism	404
12.1.6.4	Folding Through an On-pathway High-Energy Intermediate	404
	Acknowledgments	406
	References	406
12.2	Characterization of Protein Folding Barriers with Rate Equilibrium Free Energy Relationships	411
	<i>Thomas Kiefhaber, Ignacio E. Sánchez, and Annett Bachmann</i>	
12.2.1	Introduction	411
12.2.2	Rate Equilibrium Free Energy Relationships	411
12.2.2.1	Linear Rate Equilibrium Free Energy Relationships in Protein Folding	414
12.2.2.2	Properties of Protein Folding Transition States Derived from Linear REFERS	418
12.2.3	Nonlinear Rate Equilibrium Free Energy Relationships in Protein Folding	420
12.2.3.1	Self-Interaction and Cross-Interaction Parameters	420
12.2.3.2	Hammond and Anti-Hammond Behavior	424
12.2.3.3	Sequential and Parallel Transition States	425
12.2.3.4	Ground State Effects	428
12.2.4	Experimental Results on the Shape of Free Energy Barriers in Protein Folding	432
12.2.4.1	Broadness of Free Energy Barriers	432
12.2.4.2	Parallel Pathways	437
12.2.5	Folding in the Absence of Enthalpy Barriers	438
12.2.6	Conclusions and Outlook	438
	Acknowledgments	439
	References	439
13	A Guide to Measuring and Interpreting ϕ-values	445
	<i>Nicholas R. Guydosh and Alan R. Fersht</i>	
13.1	Introduction	445
13.2	Basic Concept of ϕ -Value Analysis	445
13.3	Further Interpretation of ϕ	448
13.4	Techniques	450
13.5	Conclusions	452
	References	452
14	Fast Relaxation Methods	454
	<i>Martin Gruebele</i>	
14.1	Introduction	454

14.2	Techniques	455
14.2.1	Fast Pressure-Jump Experiments	455
14.2.2	Fast Resistive Heating Experiments	456
14.2.3	Fast Laser-induced Relaxation Experiments	457
14.2.3.1	Laser Photolysis	457
14.2.3.2	Electrochemical Jumps	458
14.2.3.3	Laser-induced pH Jumps	458
14.2.3.4	Covalent Bond Dissociation	459
14.2.3.5	Chromophore Excitation	460
14.2.3.6	Laser Temperature Jumps	460
14.2.4	Multichannel Detection Techniques for Relaxation Studies	461
14.2.4.1	Small Angle X-ray Scattering or Light Scattering	462
14.2.4.2	Direct Absorption Techniques	463
14.2.4.3	Circular Dichroism and Optical Rotatory Dispersion	464
14.2.4.4	Raman and Resonance Raman Scattering	464
14.2.4.5	Intrinsic Fluorescence	465
14.2.4.6	Extrinsic Fluorescence	465
14.3	Protein Folding by Relaxation	466
14.3.1	Transition State Theory, Energy Landscapes, and Fast Folding	466
14.3.2	Viscosity Dependence of Folding Motions	470
14.3.3	Resolving Burst Phases	471
14.3.4	Fast Folding and Unfolded Proteins	472
14.3.5	Experiment and Simulation	472
14.4	Summary	474
14.5	Experimental Protocols	475
14.5.1	Design Criteria for Laser Temperature Jumps	475
14.5.2	Design Criteria for Fast Single-Shot Detection Systems	476
14.5.3	Designing Proteins for Fast Relaxation Experiments	477
14.5.4	Linear Kinetic, Nonlinear Kinetic, and Generalized Kinetic Analysis of Fast Relaxation	477
14.5.4.1	The Reaction $D \rightleftharpoons F$ in the Presence of a Barrier	477
14.5.4.2	The Reaction $2A \rightleftharpoons A_2$ in the Presence of a Barrier	478
14.5.4.3	The Reaction $D \rightleftharpoons F$ at Short Times or over Low Barriers	479
14.5.5	Relaxation Data Analysis by Linear Decomposition	480
14.5.5.1	Singular Value Decomposition (SVD)	480
14.5.5.2	χ -Analysis	481
	Acknowledgments	481
	References	482
15	Early Events in Protein Folding Explored by Rapid Mixing Methods	491
	<i>Heinrich Roder, Kosuke Maki, Ramil F. Latypov, Hong Cheng, and M. C. Ramachandra Shastry</i>	
15.1	Importance of Kinetics for Understanding Protein Folding	491
15.2	Burst-phase Signals in Stopped-flow Experiments	492
15.3	Turbulent Mixing	494

15.4	Detection Methods	495
15.4.1	Tryptophan Fluorescence	495
15.4.2	ANS Fluorescence	498
15.4.3	FRET	499
15.4.4	Continuous-flow Absorbance	501
15.4.5	Other Detection Methods used in Ultrafast Folding Studies	502
15.5	A Quenched-Flow Method for H-D Exchange Labeling Studies on the Microsecond Time Scale	502
15.6	Evidence for Accumulation of Early Folding Intermediates in Small Proteins	505
15.6.1	B1 Domain of Protein G	505
15.6.2	Ubiquitin	508
15.6.3	Cytochrome <i>c</i>	512
15.7	Significance of Early Folding Events	515
15.7.1	Barrier-limited Folding vs. Chain Diffusion	515
15.7.2	Chain Compaction: Random Collapse vs. Specific Folding	516
15.7.3	Kinetic Role of Early Folding Intermediates	517
15.7.4	Broader Implications	520
	Appendix	521
A1	Design and Calibration of Rapid Mixing Instruments	521
A1.1	Stopped-flow Equipment	521
A1.2	Continuous-flow Instrumentation	524
	Acknowledgments	528
	References	528
16	Kinetic Protein Folding Studies using NMR Spectroscopy	536
	<i>Markus Zeeb and Jochen Balbach</i>	
16.1	Introduction	536
16.2	Following Slow Protein Folding Reactions in Real Time	538
16.3	Two-dimensional Real-time NMR Spectroscopy	545
16.4	Dynamic and Spin Relaxation NMR for Quantifying Microsecond-to-Millisecond Folding Rates	550
16.5	Conclusions and Future Directions	555
16.6	Experimental Protocols	556
16.6.1	How to Record and Analyze 1D Real-time NMR Spectra	556
16.6.1.1	Acquisition	556
16.6.1.2	Processing	557
16.6.1.3	Analysis	557
16.6.1.4	Analysis of 1D Real-time Diffusion Experiments	558
16.6.2	How to Extract Folding Rates from 1D Spectra by Line Shape Analysis	559
16.6.2.1	Acquisition	560
16.6.2.2	Processing	560
16.6.2.3	Analysis	561
16.6.3	How to Extract Folding Rates from 2D Real-time NMR Spectra	562

16.6.3.1	Acquisition	563
16.6.3.2	Processing	563
16.6.3.3	Analysis	563
16.6.4	How to Analyze Heteronuclear NMR Relaxation and Exchange Data	565
16.6.4.1	Acquisition	566
16.6.4.2	Processing	567
16.6.4.3	Analysis	567
	Acknowledgments	569
	References	569

Part I, Volume 2

17	Fluorescence Resonance Energy Transfer (FRET) and Single Molecule Fluorescence Detection Studies of the Mechanism of Protein Folding and Unfolding	573
	<i>Elisha Haas</i>	
	Abbreviations	573
17.1	Introduction	573
17.2	What are the Main Aspects of the Protein Folding Problem that can be Addressed by Methods Based on FRET Measurements?	574
17.2.1	The Three Protein Folding Problems	574
17.2.1.1	The Chain Entropy Problem	574
17.2.1.2	The Function Problem: Conformational Fluctuations	575
17.3	Theoretical Background	576
17.3.1	Nonradiative Excitation Energy Transfer	576
17.3.2	What is FRET? The Singlet–Singlet Excitation Transfer	577
17.3.3	Rate of Nonradiative Excitation Energy Transfer within a Donor–Acceptor Pair	578
17.3.4	The Orientation Factor	583
17.3.5	How to Determine and Control the Value of R_0 ?	584
17.3.6	Index of Refraction n	584
17.3.7	The Donor Quantum Yield Φ_D^0	586
17.3.8	The Spectral Overlap Integral J	586
17.4	Determination of Intramolecular Distances in Protein Molecules using FRET Measurements	586
17.4.1	Single Distance between Donor and Acceptor	587
17.4.1.1	Method 1: Steady State Determination of Decrease of Donor Emission	587
17.4.1.2	Method 2: Acceptor Excitation Spectroscopy	588
17.4.2	Time-resolved Methods	588
17.4.3	Determination of E from Donor Fluorescence Decay Rates	589
17.4.4	Determination of Acceptor Fluorescence Lifetime	589
17.4.5	Determination of Intramolecular Distance Distributions	590

17.4.6	Evaluation of the Effect of Fast Conformational Fluctuations and Determination of Intramolecular Diffusion Coefficients	592
17.5	Experimental Challenges in the Implementation of FRET Folding Experiments	594
17.5.1	Optimized Design and Preparation of Labeled Protein Samples for FRET Folding Experiments	594
17.5.2	Strategies for Site-specific Double Labeling of Proteins	595
17.5.3	Preparation of Double-labeled Mutants Using Engineered Cysteine Residues (strategy 4)	596
17.5.4	Possible Pitfalls Associated with the Preparation of Labeled Protein Samples for FRET Folding Experiments	599
17.6	Experimental Aspects of Folding Studies by Distance Determination Based on FRET Measurements	600
17.6.1	Steady State Determination of Transfer Efficiency	600
17.6.1.1	Donor Emission	600
17.6.1.2	Acceptor Excitation Spectroscopy	601
17.6.2	Time-resolved Measurements	601
17.7	Data Analysis	603
17.7.1	Rigorous Error Analysis	606
17.7.2	Elimination of Systematic Errors	606
17.8	Applications of τ FRET for Characterization of Unfolded and Partially Folded Conformations of Globular Proteins under Equilibrium Conditions	607
17.8.1	Bovine Pancreatic Trypsin Inhibitor	607
17.8.2	The Loop Hypothesis	608
17.8.3	RNase A	609
17.8.4	Staphylococcal Nuclease	611
17.9	Unfolding Transition via Continuum of Native-like Forms	611
17.10	The Third Folding Problem: Domain Motions and Conformational Fluctuations of Enzyme Molecules	611
17.11	Single Molecule FRET-detected Folding Experiments	613
17.12	Principles of Applications of Single Molecule FRET Spectroscopy in Folding Studies	615
17.12.1	Design and Analysis of Single Molecule FRET Experiments	615
17.12.1.1	How is Single Molecule FRET Efficiency Determined?	615
17.12.1.2	The Challenge of Extending the Length of the Time Trajectories	617
17.12.2	Distance and Time Resolution of the Single Molecule FRET Folding Experiments	618
17.13	Folding Kinetics	619
17.13.1	Steady State and τ FRET-detected Folding Kinetics Experiments	619
17.13.2	Steady State Detection	619
17.13.3	Time-resolved FRET Detection of Rapid Folding Kinetics: the “Double Kinetics” Experiment	621
17.13.4	Multiple Probes Analysis of the Folding Transition	622
17.14	Concluding Remarks	625

	Acknowledgments	626
	References	627
18	Application of Hydrogen Exchange Kinetics to Studies of Protein Folding	634
	<i>Kaare Teilum, Birthe B. Kragelund, and Flemming M. Poulsen</i>	
18.1	Introduction	634
18.2	The Hydrogen Exchange Reaction	638
18.2.1	Calculating the Intrinsic Hydrogen Exchange Rate Constant, k_{int}	638
18.3	Protein Dynamics by Hydrogen Exchange in Native and Denaturing Conditions	641
18.3.1	Mechanisms of Exchange	642
18.3.2	Local Opening and Closing Rates from Hydrogen Exchange Kinetics	642
18.3.2.1	The General Amide Exchange Rate Expression – the Linderstrøm-Lang Equation	643
18.3.2.2	Limits to the General Rate Expression – EX1 and EX2	644
18.3.2.3	The Range between the EX1 and EX2 Limits	646
18.3.2.4	Identification of Exchange Limit	646
18.3.2.5	Global Opening and Closing Rates and Protein Folding	647
18.3.3	The “Native State Hydrogen Exchange” Strategy	648
18.3.3.1	Localization of Partially Unfolded States, PUFs	650
18.4	Hydrogen Exchange as a Structural Probe in Kinetic Folding Experiments	651
18.4.1	Protein Folding/Hydrogen Exchange Competition	652
18.4.2	Hydrogen Exchange Pulse Labeling	656
18.4.3	Protection Factors in Folding Intermediates	657
18.4.4	Kinetic Intermediate Structures Characterized by Hydrogen Exchange	659
18.5	Experimental Protocols	661
18.5.1	How to Determine Hydrogen Exchange Kinetics at Equilibrium	661
18.5.1.1	Equilibrium Hydrogen Exchange Experiments	661
18.5.1.2	Determination of Segmental Opening and Closing Rates, k_{op} and k_{cl}	662
18.5.1.3	Determination of ΔG_{fluc} , m , and $\Delta G^{\circ}_{\text{unf}}$	662
18.5.2	Planning a Hydrogen Exchange Folding Experiment	662
18.5.2.1	Determine a Combination of t_{pulse} and pH_{pulse}	662
18.5.2.2	Setup Quench Flow Apparatus	662
18.5.2.3	Prepare Deuterated Protein and Chemicals	663
18.5.2.4	Prepare Buffers and Unfolded Protein	663
18.5.2.5	Check pH in the Mixing Steps	664
18.5.2.6	Sample Mixing and Preparation	664
18.5.3	Data Analysis	664
	Acknowledgments	665
	References	665

19	Studying Protein Folding and Aggregation by Laser Light Scattering	673
	<i>Klaus Gast and Andreas J. Modler</i>	
19.1	Introduction	673
19.2	Basic Principles of Laser Light Scattering	674
19.2.1	Light Scattering by Macromolecular Solutions	674
19.2.2	Molecular Parameters Obtained from Static Light Scattering (SLS)	676
19.2.3	Molecular Parameters Obtained from Dynamic Light Scattering (DLS)	678
19.2.4	Advantages of Combined SLS and DLS Experiments	680
19.3	Laser Light Scattering of Proteins in Different Conformational States – Equilibrium Folding/Unfolding Transitions	680
19.3.1	General Considerations, Hydrodynamic Dimensions in the Natively Folded State	680
19.3.2	Changes in the Hydrodynamic Dimensions during Heat-induced Unfolding	682
19.3.3	Changes in the Hydrodynamic Dimensions upon Cold Denaturation	683
19.3.4	Denaturant-induced Changes of the Hydrodynamic Dimensions	684
19.3.5	Acid-induced Changes of the Hydrodynamic Dimensions	685
19.3.6	Dimensions in Partially Folded States – Molten Globules and Fluoroalcohol-induced States	686
19.3.7	Comparison of the Dimensions of Proteins in Different Conformational States	687
19.3.8	Scaling Laws for the Native and Highly Unfolded States, Hydrodynamic Modeling	687
19.4	Studying Folding Kinetics by Laser Light Scattering	689
19.4.1	General Considerations, Attainable Time Regions	689
19.4.2	Hydrodynamic Dimensions of the Kinetic Molten Globule of Bovine α -Lactalbumin	690
19.4.3	RNase A is Only Weakly Collapsed During the Burst Phase of Folding	691
19.5	Misfolding and Aggregation Studied by Laser Light Scattering	692
19.5.1	Overview: Some Typical Light Scattering Studies of Protein Aggregation	692
19.5.2	Studying Misfolding and Amyloid Formation by Laser Light Scattering	693
19.5.2.1	Overview: Initial States, Critical Oligomers, Protofibrils, Fibrils	693
19.5.2.2	Aggregation Kinetics of $A\beta$ Peptides	694
19.5.2.3	Kinetics of Oligomer and Fibril Formation of PGK and Recombinant Hamster Prion Protein	695
19.5.2.4	Mechanisms of Misfolding and Misassembly, Some General Remarks	698
19.6	Experimental Protocols	698
19.6.1	Laser Light Scattering Instrumentation	698

19.6.1.1	Basic Experimental Set-up, General Requirements	698
19.6.1.2	Supplementary Measurements and Useful Options	700
19.6.1.3	Commercially Available Light Scattering Instrumentation	701
19.6.2	Experimental Protocols for the Determination of Molecular Mass and Stokes Radius of a Protein in a Particular Conformational State	701
	Protocol 1	702
	Protocol 2	704
	Acknowledgments	704
	References	704

20 Conformational Properties of Unfolded Proteins 710

Patrick J. Fleming and George D. Rose

20.1	Introduction	710
20.1.1	Unfolded vs. Denatured Proteins	710
20.2	Early History	711
20.3	The Random Coil	712
20.3.1	The Random Coil – Theory	713
20.3.1.1	The Random Coil Model Prompts Three Questions	716
20.3.1.2	The Folding Funnel	716
20.3.1.3	Transition State Theory	717
20.3.1.4	Other Examples	717
20.3.1.5	Implicit Assumptions from the Random Coil Model	718
20.3.2	The Random Coil – Experiment	718
20.3.2.1	Intrinsic Viscosity	719
20.3.2.2	SAXS and SANS	720
20.4	Questions about the Random Coil Model	721
20.4.1	Questions from Theory	722
20.4.1.1	The Flory Isolated-pair Hypothesis	722
20.4.1.2	Structure vs. Energy Duality	724
20.4.1.3	The “Rediscovery” of Polyproline II Conformation	724
20.4.1.4	P _{II} in Unfolded Peptides and Proteins	726
20.4.2	Questions from Experiment	727
20.4.2.1	Residual Structure in Denatured Proteins and Peptides	727
20.4.3	The Reconciliation Problem	728
20.4.4	Organization in the Unfolded State – the Entropic Conjecture	728
20.4.4.1	Steric Restrictions beyond the Dipeptide	729
20.5	Future Directions	730
	Acknowledgments	731
	References	731

21 Conformation and Dynamics of Nonnative States of Proteins studied by NMR Spectroscopy 737

Julia Wirmer, Christian Schlörb, and Harald Schwalbe

21.1	Introduction	737
21.1.1	Structural Diversity of Polypeptide Chains	737

21.1.2	Intrinsically Unstructured and Natively Unfolded Proteins	739
21.2	Prerequisites: NMR Resonance Assignment	740
21.3	NMR Parameters	744
21.3.1	Chemical shifts δ	745
21.3.1.1	Conformational Dependence of Chemical Shifts	745
21.3.1.2	Interpretation of Chemical Shifts in the Presence of Conformational Averaging	746
21.3.2	J Coupling Constants	748
21.3.2.1	Conformational Dependence of J Coupling Constants	748
21.3.2.2	Interpretation of J Coupling Constants in the Presence of Conformational Averaging	750
21.3.3	Relaxation: Homonuclear NOEs	750
21.3.3.1	Distance Dependence of Homonuclear NOEs	750
21.3.3.2	Interpretation of Homonuclear NOEs in the Presence of Conformational Averaging	754
21.3.4	Heteronuclear Relaxation (^{15}N R_1 , R_2 , hetNOE)	757
21.3.4.1	Correlation Time Dependence of Heteronuclear Relaxation Parameters	757
21.3.4.2	Dependence on Internal Motions of Heteronuclear Relaxation Parameters	759
21.3.5	Residual Dipolar Couplings	760
21.3.5.1	Conformational Dependence of Residual Dipolar Couplings	760
21.3.5.2	Interpretation of Residual Dipolar Couplings in the Presence of Conformational Averaging	763
21.3.6	Diffusion	765
21.3.7	Paramagnetic Spin Labels	766
21.3.8	H/D Exchange	767
21.3.9	Photo-CIDNP	767
21.4	Model for the Random Coil State of a Protein	768
21.5	Nonnative States of Proteins: Examples from Lysozyme, α -Lactalbumin, and Ubiquitin	771
21.5.1	Backbone Conformation	772
21.5.1.1	Interpretation of Chemical Shifts	772
21.5.1.2	Interpretation of NOEs	774
21.5.1.3	Interpretation of J Coupling Constants	780
21.5.2	Side-chain Conformation	784
21.5.2.1	Interpretation of J Coupling Constants	784
21.5.3	Backbone Dynamics	786
21.5.3.1	Interpretation of ^{15}N Relaxation Rates	786
21.6	Summary and Outlook	793
	Acknowledgments	794
	References	794
22	Dynamics of Unfolded Polypeptide Chains	809
	<i>Beat Fierz and Thomas Kiefhaber</i>	
22.1	Introduction	809

22.2	Equilibrium Properties of Chain Molecules	809
22.2.1	The Freely Jointed Chain	810
22.2.2	Chain Stiffness	810
22.2.3	Polypeptide Chains	811
22.2.4	Excluded Volume Effects	812
22.3	Theory of Polymer Dynamics	813
22.3.1	The Langevin Equation	813
22.3.2	Rouse Model and Zimm Model	814
22.3.3	Dynamics of Loop Closure and the Szabo-Schulten-Schulten Theory	815
22.4	Experimental Studies on the Dynamics in Unfolded Polypeptide Chains	816
22.4.1	Experimental Systems for the Study of Intrachain Diffusion	816
22.4.1.1	Early Experimental Studies	816
22.4.1.2	Triplet Transfer and Triplet Quenching Studies	821
22.4.1.3	Fluorescence Quenching	825
22.4.2	Experimental Results on Dynamic Properties of Unfolded Polypeptide Chains	825
22.4.2.1	Kinetics of Intrachain Diffusion	826
22.4.2.2	Effect of Loop Size on the Dynamics in Flexible Polypeptide Chains	826
22.4.2.3	Effect of Amino Acid Sequence on Chain Dynamics	829
22.4.2.4	Effect of the Solvent on Intrachain Diffusion	831
22.4.2.5	Effect of Solvent Viscosity on Intrachain Diffusion	833
22.4.2.6	End-to-end Diffusion vs. Intrachain Diffusion	834
22.4.2.7	Chain Diffusion in Natural Protein Sequences	834
22.5	Implications for Protein Folding Kinetics	837
22.5.1	Rate of Contact Formation during the Earliest Steps in Protein Folding	837
22.5.2	The Speed Limit of Protein Folding vs. the Pre-exponential Factor	839
22.5.3	Contributions of Chain Dynamics to Rate- and Equilibrium Constants for Protein Folding Reactions	840
22.6	Conclusions and Outlook	844
22.7	Experimental Protocols and Instrumentation	844
22.7.1	Properties of the Electron Transfer Probes and Treatment of the Transfer Kinetics	845
22.7.2	Test for Diffusion-controlled Reactions	847
22.7.2.1	Determination of Bimolecular Quenching or Transfer Rate Constants	847
22.7.2.2	Testing the Viscosity Dependence	848
22.7.2.3	Determination of Activation Energy	848
22.7.3	Instrumentation	849
	Acknowledgments	849
	References	849

23	Equilibrium and Kinetically Observed Molten Globule States	856
	<i>Kosuke Maki, Kiyoto Kamagata, and Kunihiro Kuwajima</i>	
23.1	Introduction	856
23.2	Equilibrium Molten Globule State	858
23.2.1	Structural Characteristics of the Molten Globule State	858
23.2.2	Typical Examples of the Equilibrium Molten Globule State	859
23.2.3	Thermodynamic Properties of the Molten Globule State	860
23.3	The Kinetically Observed Molten Globule State	862
23.3.1	Observation and Identification of the Molten Globule State in Kinetic Refolding	862
23.3.2	Kinetics of Formation of the Early Folding Intermediates	863
23.3.3	Late Folding Intermediates and Structural Diversity	864
23.3.4	Evidence for the On-pathway Folding Intermediate	865
23.4	Two-stage Hierarchical Folding Funnel	866
23.5	Unification of the Folding Mechanism between Non-two-state and Two-state Proteins	867
23.5.1	Statistical Analysis of the Folding Data of Non-two-state and Two-state Proteins	868
23.5.2	A Unified Mechanism of Protein Folding: Hierarchy	870
23.5.3	Hidden Folding Intermediates in Two-state Proteins	871
23.6	Practical Aspects of the Experimental Study of Molten Globules	872
23.6.1	Observation of the Equilibrium Molten Globule State	872
23.6.1.1	Two-state Unfolding Transition	872
23.6.1.2	Multi-state (Three-state) Unfolding Transition	874
23.6.2	Burst-phase Intermediate Accumulated during the Dead Time of Refolding Kinetics	876
23.6.3	Testing the Identity of the Molten Globule State with the Burst-Phase Intermediate	877
	References	879
24	Alcohol- and Salt-induced Partially Folded Intermediates	884
	<i>Daizo Hamada and Yuji Goto</i>	
24.1	Introduction	884
24.2	Alcohol-induced Intermediates of Proteins and Peptides	886
24.2.1	Formation of Secondary Structures by Alcohols	888
24.2.2	Alcohol-induced Denaturation of Proteins	888
24.2.3	Formation of Compact Molten Globule States	889
24.2.4	Example: β -Lactoglobulin	890
24.3	Mechanism of Alcohol-induced Conformational Change	893
24.4	Effects of Alcohols on Folding Kinetics	896
24.5	Salt-induced Formation of the Intermediate States	899
24.5.1	Acid-denatured Proteins	899
24.5.2	Acid-induced Unfolding and Refolding Transitions	900
24.6	Mechanism of Salt-induced Conformational Change	904
24.7	Generality of the Salt Effects	906

24.8	Conclusion	907
	References	908
25	Prolyl Isomerization in Protein Folding	916
	<i>Franz Schmid</i>	
25.1	Introduction	916
25.2	Prolyl Peptide Bonds	917
25.3	Prolyl Isomerizations as Rate-determining Steps of Protein Folding	918
25.3.1	The Discovery of Fast and Slow Refolding Species	918
25.3.2	Detection of Proline-limited Folding Processes	919
25.3.3	Proline-limited Folding Reactions	921
25.3.4	Interrelation between Prolyl Isomerization and Conformational Folding	923
25.4	Examples of Proline-limited Folding Reactions	924
25.4.1	Ribonuclease A	924
25.4.2	Ribonuclease T1	926
25.4.3	The Structure of a Folding Intermediate with an Incorrect Prolyl Isomer	928
25.5	Native-state Prolyl Isomerizations	929
25.6	Nonprolyl Isomerizations in Protein Folding	930
25.7	Catalysis of Protein Folding by Prolyl Isomerases	932
25.7.1	Prolyl Isomerases as Tools for Identifying Proline-limited Folding Steps	932
25.7.2	Specificity of Prolyl Isomerases	933
25.7.3	The Trigger Factor	934
25.7.4	Catalysis of Prolyl Isomerization During de novo Protein Folding	935
25.8	Concluding Remarks	936
25.9	Experimental Protocols	936
25.9.1	Slow Refolding Assays ("Double Jumps") to Measure Prolyl Isomerizations in an Unfolded Protein	936
25.9.1.1	Guidelines for the Design of Double Jump Experiments	937
25.9.1.2	Formation of U _S Species after Unfolding of RNase A	938
25.9.2	Slow Unfolding Assays for Detecting and Measuring Prolyl Isomerizations in Refolding	938
25.9.2.1	Practical Considerations	939
25.9.2.2	Kinetics of the Formation of Fully Folded IIHY-G3P* Molecules	939
	References	939
26	Folding and Disulfide Formation	946
	<i>Margherita Ruoppolo, Piero Pucci, and Gennaro Marino</i>	
26.1	Chemistry of the Disulfide Bond	946
26.2	Trapping Protein Disulfides	947
26.3	Mass Spectrometric Analysis of Folding Intermediates	948
26.4	Mechanism(s) of Oxidative Folding so Far – Early and Late Folding Steps	949

26.5	Emerging Concepts from Mass Spectrometric Studies	950
26.5.1	Three-fingered Toxins	951
26.5.2	RNase A	953
26.5.3	Antibody Fragments	955
26.5.4	Human Nerve Growth Factor	956
26.6	Unanswered Questions	956
26.7	Concluding Remarks	957
26.8	Experimental Protocols	957
26.8.1	How to Prepare Folding Solutions	957
26.8.2	How to Carry Out Folding Reactions	958
26.8.3	How to Choose the Best Mass Spectrometric Equipment for Your Study	959
26.8.4	How to Perform Electrospray (ES)MS Analysis	959
26.8.5	How to Perform Matrix-assisted Laser Desorption Ionization (MALDI) MS Analysis	960
	References	961
27	Concurrent Association and Folding of Small Oligomeric Proteins	965
	<i>Hans Rudolf Bosshard</i>	
27.1	Introduction	965
27.2	Experimental Methods Used to Follow the Folding of Oligomeric Proteins	966
27.2.1	Equilibrium Methods	966
27.2.2	Kinetic Methods	968
27.3	Dimeric Proteins	969
27.3.1	Two-state Folding of Dimeric Proteins	970
27.3.1.1	Examples of Dimeric Proteins Obeying Two-state Folding	971
27.3.2	Folding of Dimeric Proteins through Intermediate States	978
27.4	Trimeric and Tetrameric Proteins	983
27.5	Concluding Remarks	986
	Appendix – Concurrent Association and Folding of Small Oligomeric Proteins	987
A1	Equilibrium Constants for Two-state Folding	988
A1.1	Homooligomeric Protein	988
A1.2	Heterooligomeric Protein	989
A2	Calculation of Thermodynamic Parameters from Equilibrium Constants	990
A2.1	Basic Thermodynamic Relationships	990
A2.2	Linear Extrapolation of Denaturant Unfolding Curves of Two-state Reaction	990
A2.3	Calculation of the van't Hoff Enthalpy Change from Thermal Unfolding Data	990
A2.4	Calculation of the van't Hoff Enthalpy Change from the Concentration-dependence of T_m	991
A2.5	Extrapolation of Thermodynamic Parameters to Different Temperatures: Gibbs-Helmholtz Equation	991

A3	Kinetics of Reversible Two-state Folding and Unfolding: Integrated Rate Equations	992
A3.1	Two-state Folding of Dimeric Protein	992
A3.2	Two-state Unfolding of Dimeric Protein	992
A3.3	Reversible Two-state Folding and Unfolding	993
A3.3.1	Homodimeric protein	993
A3.3.2	Heterodimeric protein	993
A4	Kinetics of Reversible Two-state Folding: Relaxation after Disturbance of a Pre-existing Equilibrium (Method of Bernasconi)	994
	Acknowledgments	995
	References	995
28	Folding of Membrane Proteins	998
	<i>Lukas K. Tamm and Heedeok Hong</i>	
28.1	Introduction	998
28.2	Thermodynamics of Residue Partitioning into Lipid Bilayers	1000
28.3	Stability of β -Barrel Proteins	1001
28.4	Stability of Helical Membrane Proteins	1009
28.5	Helix and Other Lateral Interactions in Membrane Proteins	1010
28.6	The Membrane Interface as an Important Contributor to Membrane Protein Folding	1012
28.7	Membrane Toxins as Models for Helical Membrane Protein Insertion	1013
28.8	Mechanisms of β -Barrel Membrane Protein Folding	1015
28.9	Experimental Protocols	1016
28.9.1	SDS Gel Shift Assay for Heat-modifiable Membrane Proteins	1016
28.9.1.1	Reversible Folding and Unfolding Protocol Using OmpA as an Example	1016
28.9.2	Tryptophan Fluorescence and Time-resolved Distance Determination by Tryptophan Fluorescence Quenching	1018
28.9.2.1	TDFQ Protocol for Monitoring the Translocation of Tryptophans across Membranes	1019
28.9.3	Circular Dichroism Spectroscopy	1020
28.9.4	Fourier Transform Infrared Spectroscopy	1022
28.9.4.1	Protocol for Obtaining Conformation and Orientation of Membrane Proteins and Peptides by Polarized ATR-FTIR Spectroscopy	1023
	Acknowledgments	1025
	References	1025
29	Protein Folding Catalysis by Pro-domains	1032
	<i>Philip N. Bryan</i>	
29.1	Introduction	1032
29.2	Bimolecular Folding Mechanisms	1033
29.3	Structures of Reactants and Products	1033
29.3.1	Structure of Free SBT	1033

29.3.2	Structure of SBT/Pro-domain Complex	1036
29.3.3	Structure of Free ALP	1037
29.3.4	Structure of the ALP/Pro-domain Complex	1037
29.4	Stability of the Mature Protease	1039
29.4.1	Stability of ALP	1039
29.4.2	Stability of Subtilisin	1040
29.5	Analysis of Pro-domain Binding to the Folded Protease	1042
29.6	Analysis of Folding Steps	1043
29.7	Why are Pro-domains Required for Folding?	1046
29.8	What is the Origin of High Cooperativity?	1047
29.9	How Does the Pro-domain Accelerate Folding?	1048
29.10	Are High Kinetic Stability and Facile Folding Mutually Exclusive?	1049
29.11	Experimental Protocols for Studying SBT Folding	1049
29.11.1	Fermentation and Purification of Active Subtilisin	1049
29.11.2	Fermentation and Purification of Facile-folding Ala221 Subtilisin from <i>E. coli</i>	1050
29.11.3	Mutagenesis and Protein Expression of Pro-domain Mutants	1051
29.11.4	Purification of Pro-domain	1052
29.11.5	Kinetics of Pro-domain Binding to Native SBT	1052
29.11.6	Kinetic Analysis of Pro-domain Facilitated Subtilisin Folding	1052
29.11.6.1	Single Mixing	1052
29.11.6.2	Double Jump: Renaturation–Denaturation	1053
29.11.6.3	Double Jump: Denaturation–Renaturation	1053
29.11.6.4	Triple Jump: Denaturation–Renaturation–Denaturation	1054
	References	1054
30	The Thermodynamics and Kinetics of Collagen Folding	1059
	<i>Hans Peter Bächinger and Jürgen Engel</i>	
30.1	Introduction	1059
30.1.1	The Collagen Family	1059
30.1.2	Biosynthesis of Collagens	1060
30.1.3	The Triple Helical Domain in Collagens and Other Proteins	1061
30.1.4	N- and C-Propeptide, Telopeptides, Flanking Coiled-Coil Domains	1061
30.1.5	Why is the Folding of the Triple Helix of Interest?	1061
30.2	Thermodynamics of Collagen Folding	1062
30.2.1	Stability of the Triple Helix	1062
30.2.2	The Role of Posttranslational Modifications	1063
30.2.3	Energies Involved in the Stability of the Triple Helix	1063
30.2.4	Model Peptides Forming the Collagen Triple Helix	1066
30.2.4.1	Type of Peptides	1066
30.2.4.2	The All-or-none Transition of Short Model Peptides	1066
30.2.4.3	Thermodynamic Parameters for Different Model Systems	1069
30.2.4.4	Contribution of Different Tripeptide Units to Stability	1075

30.2.4.5	Crystal and NMR Structures of Triple Helices	1076
30.2.4.6	Conformation of the Randomly Coiled Chains	1077
30.2.4.7	Model Studies with Isomers of Hydroxyproline and Fluoroproline	1078
30.2.4.8	Cis \rightleftharpoons trans Equilibria of Peptide Bonds	1079
30.2.4.9	Interpretations of Stabilities on a Molecular Level	1080
30.3	Kinetics of Triple Helix Formation	1081
30.3.1	Properties of Collagen Triple Helices that Influence Kinetics	1081
30.3.2	Folding of Triple Helices from Single Chains	1082
30.3.2.1	Early Work	1082
30.3.2.2	Concentration Dependence of the Folding of (PPG) ₁₀ and (POG) ₁₀	1082
30.3.2.3	Model Mechanism of the Folding Kinetics	1085
30.3.2.4	Rate Constants of Nucleation and Propagation	1087
30.3.2.5	Host-guest Peptides and an Alternative Kinetics Model	1088
30.3.3	Triple Helix Formation from Linked Chains	1089
30.3.3.1	The Short N-terminal Triple Helix of Collagen III in Fragment Col1-3	1089
30.3.3.2	Folding of the Central Long Triple Helix of Collagen III	1090
30.3.3.3	The Zipper Model	1092
30.3.4	Designed Collagen Models with Chains Connected by a Disulfide Knot or by Trimerizing Domains	1097
30.3.4.1	Disulfide-linked Model Peptides	1097
30.3.4.2	Model Peptides Linked by a Foldon Domain	1098
30.3.4.3	Collagen Triple Helix Formation can be Nucleated at either End	1098
30.3.4.4	Hysteresis of Triple Helix Formation	1099
30.3.5	Influence of <i>cis-trans</i> Isomerase and Chaperones	1100
30.3.6	Mutations in Collagen Triple Helices Affect Proper Folding	1101
	References	1101
31	Unfolding Induced by Mechanical Force	1111
	<i>Jane Clarke and Phil M. Williams</i>	
31.1	Introduction	1111
31.2	Experimental Basics	1112
31.2.1	Instrumentation	1112
31.2.2	Sample Preparation	1113
31.2.3	Collecting Data	1114
31.2.4	Anatomy of a Force Trace	1115
31.2.5	Detecting Intermediates in a Force Trace	1115
31.2.6	Analyzing the Force Trace	1116
31.3	Analysis of Force Data	1117
31.3.1	Basic Theory behind Dynamic Force Spectroscopy	1117
31.3.2	The Ramp of Force Experiment	1119
31.3.3	The Golden Equation of DFS	1121
31.3.4	Nonlinear Loading	1122

31.3.4.1	The Worm-line Chain (WLC)	1123
31.3.5	Experiments under Constant Force	1124
31.3.6	Effect of Tandem Repeats on Kinetics	1125
31.3.7	Determining the Modal Force	1126
31.3.8	Comparing Behavior	1127
31.3.9	Fitting the Data	1127
31.4	Use of Complementary Techniques	1129
31.4.1	Protein Engineering	1130
31.4.1.1	Choosing Mutants	1130
31.4.1.2	Determining $\Delta\Delta G_{D-N}$	1131
31.4.1.3	Determining $\Delta\Delta G_{TS-N}$	1131
31.4.1.4	Interpreting the Φ -values	1132
31.4.2	Computer Simulation	1133
31.5	Titin I27: A Case Study	1134
31.5.1	The Protein System	1134
31.5.2	The Unfolding Intermediate	1135
31.5.3	The Transition State	1136
31.5.4	The Relationship Between the Native and Transition States	1137
31.5.5	The Energy Landscape under Force	1139
31.6	Conclusions – the Future	1139
	References	1139
32	Molecular Dynamics Simulations to Study Protein Folding and Unfolding	1143
	<i>Amedeo Caflisch and Emanuele Paci</i>	
32.1	Introduction	1143
32.2	Molecular Dynamics Simulations of Peptides and Proteins	1144
32.2.1	Folding of Structured Peptides	1144
32.2.1.1	Reversible Folding and Free Energy Surfaces	1144
32.2.1.2	Non-Arrhenius Temperature Dependence of the Folding Rate	1147
32.2.1.3	Denatured State and Levinthal Paradox	1148
32.2.1.4	Folding Events of Trp-cage	1149
32.2.2	Unfolding Simulations of Proteins	1150
32.2.2.1	High-temperature Simulations	1150
32.2.2.2	Biased Unfolding	1150
32.2.2.3	Forced Unfolding	1151
32.2.3	Determination of the Transition State Ensemble	1153
32.3	MD Techniques and Protocols	1155
32.3.1	Techniques to Improve Sampling	1155
32.3.1.1	Replica Exchange Molecular Dynamics	1155
32.3.1.2	Methods Based on Path Sampling	1157
32.3.2	MD with Restraints	1157
32.3.3	Distributed Computing Approach	1158
32.3.4	Implicit Solvent Models versus Explicit Water	1160
32.4	Conclusion	1162
	References	1162

33	Molecular Dynamics Simulations of Proteins and Peptides: Problems, Achievements, and Perspectives	1170
	<i>Paul Tavan, Heiko Carstens, and Gerald Mathias</i>	
33.1	Introduction	1170
33.2	Basic Physics of Protein Structure and Dynamics	1171
33.2.1	Protein Electrostatics	1172
33.2.2	Relaxation Times and Spatial Scales	1172
33.2.3	Solvent Environment	1173
33.2.4	Water	1174
33.2.5	Polarizability of the Peptide Groups and of Other Protein Components	1175
33.3	State of the Art	1177
33.3.1	Control of Thermodynamic Conditions	1177
33.3.2	Long-range Electrostatics	1177
33.3.3	Polarizability	1179
33.3.4	Higher Multipole Moments of the Molecular Components	1180
33.3.5	MM Models of Water	1181
33.3.6	Complexity of Protein–Solvent Systems and Consequences for MM-MD	1182
33.3.7	What about Successes of MD Methods?	1182
33.3.8	Accessible Time Scales and Accuracy Issues	1184
33.3.9	Continuum Solvent Models	1185
33.3.10	Are there Further Problems beyond Electrostatics and Structure Prediction?	1187
33.4	Conformational Dynamics of a Light-switchable Model Peptide	1187
33.4.1	Computational Methods	1188
33.4.2	Results and Discussion	1190
	Summary	1194
	Acknowledgments	1194
	References	1194

Part II, Volume 1

Contributors of Part II LVIII

1	Paradigm Changes from “Unboiling an Egg” to “Synthesizing a Rabbit”	3
	<i>Rainer Jaenicke</i>	
1.1	Protein Structure, Stability, and Self-organization	3
1.2	Autonomous and Assisted Folding and Association	6
1.3	Native, Intermediate, and Denatured States	11
1.4	Folding and Merging of Domains – Association of Subunits	13
1.5	Limits of Reconstitution	19
1.6	In Vitro Denaturation-Renaturation vs. Folding in Vivo	21

1.7	Perspectives	24
	Acknowledgements	26
	References	26
2	Folding and Association of Multi-domain and Oligomeric Proteins	32
	<i>Hauke Lilie and Robert Seckler</i>	
2.1	Introduction	32
2.2	Folding of Multi-domain Proteins	33
2.2.1	Domain Architecture	33
2.2.2	γ -Crystallin as a Model for a Two-domain Protein	35
2.2.3	The Giant Protein Titin	39
2.3	Folding and Association of Oligomeric Proteins	41
2.3.1	Why Oligomers?	41
2.3.2	Inter-subunit Interfaces	42
2.3.3	Domain Swapping	44
2.3.4	Stability of Oligomeric Proteins	45
2.3.5	Methods Probing Folding/Association	47
2.3.5.1	Chemical Cross-linking	47
2.3.5.2	Analytical Gel Filtration Chromatography	47
2.3.5.3	Scattering Methods	48
2.3.5.4	Fluorescence Resonance Energy Transfer	48
2.3.5.5	Hybrid Formation	48
2.3.6	Kinetics of Folding and Association	49
2.3.6.1	General Considerations	49
2.3.6.2	Reconstitution Intermediates	50
2.3.6.3	Rates of Association	52
2.3.6.4	Homo- Versus Heterodimerization	52
2.4	Renaturation versus Aggregation	54
2.5	Case Studies on Protein Folding and Association	54
2.5.1	Antibody Fragments	54
2.5.2	Trimeric Tail Spike Protein of Bacteriophage P22	59
2.6	Experimental Protocols	62
	References	65
3	Studying Protein Folding in Vivo	73
	<i>I. Marije Liscajet, Bertrand Kleizen, and Ineke Braakman</i>	
3.1	Introduction	73
3.2	General Features in Folding Proteins Amenable to in Vivo Study	73
3.2.1	Increasing Compactness	76
3.2.2	Decreasing Accessibility to Different Reagents	76
3.2.3	Changes in Conformation	77
3.2.4	Assistance During Folding	78
3.3	Location-specific Features in Protein Folding	79
3.3.1	Translocation and Signal Peptide Cleavage	79
3.3.2	Glycosylation	80

3.3.3	Disulfide Bond Formation in the ER	81
3.3.4	Degradation	82
3.3.5	Transport from ER to Golgi and Plasma Membrane	83
3.4	How to Manipulate Protein Folding	84
3.4.1	Pharmacological Intervention (Low-molecular-weight Reagents)	84
3.4.1.1	Reducing and Oxidizing Agents	84
3.4.1.2	Calcium Depletion	84
3.4.1.3	ATP Depletion	85
3.4.1.4	Cross-linking	85
3.4.1.5	Glycosylation Inhibitors	85
3.4.2	Genetic Modifications (High-molecular-weight Manipulations)	86
3.4.2.1	Substrate Protein Mutants	86
3.4.2.2	Changing the Concentration or Activity of Folding Enzymes and Chaperones	87
3.5	Experimental Protocols	88
3.5.1	Protein-labeling Protocols	88
3.5.1.1	Basic Protocol Pulse Chase: Adherent Cells	88
3.5.1.2	Pulse Chase in Suspension Cells	91
3.5.2	(Co)-immunoprecipitation and Accessory Protocols	93
3.5.2.1	Immunoprecipitation	93
3.5.2.2	Co-precipitation with Calnexin ([84]; adapted from Ou et al. [85])	94
3.5.2.3	Co-immunoprecipitation with Other Chaperones	95
3.5.2.4	Protease Resistance	95
3.5.2.5	Endo H Resistance	96
3.5.2.6	Cell Surface Expression Tested by Protease	96
3.5.3	SDS-PAGE [13]	97
	Acknowledgements	98
	References	98
4	Characterization of ATPase Cycles of Molecular Chaperones by Fluorescence and Transient Kinetic Methods	105
	<i>Sandra Schlee and Jochen Reinstein</i>	
4.1	Introduction	105
4.1.1	Characterization of ATPase Cycles of Energy-transducing Systems	105
4.1.2	The Use of Fluorescent Nucleotide Analogues	106
4.1.2.1	Fluorescent Modifications of Nucleotides	106
4.1.2.2	How to Find a Suitable Analogue for a Specific Protein	108
4.2	Characterization of ATPase Cycles of Molecular Chaperones	109
4.2.1	Biased View	109
4.2.2	The ATPase Cycle of DnaK	109
4.2.3	The ATPase Cycle of the Chaperone Hsp90	109
4.2.4	The ATPase Cycle of the Chaperone ClpB	111
4.2.4.1	ClpB, an Oligomeric ATPase With Two AAA Modules Per Protomer	111

4.2.4.2	Nucleotide-binding Properties of NBD1 and NBD2	111
4.2.4.3	Cooperativity of ATP Hydrolysis and Interdomain Communication	114
4.3	Experimental Protocols	116
4.3.1	Synthesis of Fluorescent Nucleotide Analogues	116
4.3.1.1	Synthesis and Characterization of (P _{β})MABA-ADP and (P _{γ})MABA-ATP	116
4.3.1.2	Synthesis and Characterization of N8-MABA Nucleotides	119
4.3.1.3	Synthesis of MANT Nucleotides	120
4.3.2	Preparation of Nucleotides and Proteins	121
4.3.2.1	Assessment of Quality of Nucleotide Stock Solution	121
4.3.2.2	Determination of the Nucleotide Content of Proteins	122
4.3.2.3	Nucleotide Depletion Methods	123
4.3.3	Steady-state ATPase Assays	124
4.3.3.1	Coupled Enzymatic Assay	124
4.3.3.2	Assays Based on [α - ³² P]-ATP and TLC	125
4.3.3.3	Assays Based on Released P _i	125
4.3.4	Single-turnover ATPase Assays	126
4.3.4.1	Manual Mixing Procedures	126
4.3.4.2	Quenched Flow	127
4.3.5	Nucleotide-binding Measurements	127
4.3.5.1	Isothermal Titration Calorimetry	127
4.3.5.2	Equilibrium Dialysis	129
4.3.5.3	Filter Binding	129
4.3.5.4	Equilibrium Fluorescence Titration	130
4.3.5.5	Competition Experiments	132
4.3.6	Analytical Solutions of Equilibrium Systems	133
4.3.6.1	Quadratic Equation	133
4.3.6.2	Cubic Equation	134
4.3.6.3	Iterative Solutions	138
4.3.7	Time-resolved Binding Measurements	141
4.3.7.1	Introduction	141
4.3.7.2	One-step Irreversible Process	142
4.3.7.3	One-step Reversible Process	143
4.3.7.4	Reversible Second Order Reduced to Pseudo-first Order	144
4.3.7.5	Two Simultaneous Irreversible Pathways – Partitioning	146
4.3.7.6	Two-step Consecutive (Sequential) Reaction	148
4.3.7.7	Two-step Binding Reactions	150
	References	152
5	Analysis of Chaperone Function in Vitro	162
	<i>Johannes Buchner and Stefan Walter</i>	
5.1	Introduction	162
5.2	Basic Functional Principles of Molecular Chaperones	164
5.2.1	Recognition of Nonnative Proteins	166

5.2.2	Induction of Conformational Changes in the Substrate	167
5.2.3	Energy Consumption and Regulation of Chaperone Function	169
5.3	Limits and Extensions of the Chaperone Concept	170
5.3.1	Co-chaperones	171
5.3.2	Specific Chaperones	171
5.4	Working with Molecular Chaperones	172
5.4.1	Natural versus Artificial Substrate Proteins	172
5.4.2	Stability of Chaperones	172
5.5	Assays to Assess and Characterize Chaperone Function	174
5.5.1	Generating Nonnative Conformations of Proteins	174
5.5.2	Aggregation Assays	174
5.5.3	Detection of Complexes Between Chaperone and Substrate	175
5.5.4	Refolding of Denatured Substrates	175
5.5.5	ATPase Activity and Effect of Substrate and Cofactors	176
5.6	Experimental Protocols	176
5.6.1	General Considerations	176
5.6.1.1	Analysis of Chaperone Stability	176
5.6.1.2	Generation of Nonnative Proteins	177
5.6.1.3	Model Substrates for Chaperone Assays	177
5.6.2	Suppression of Aggregation	179
5.6.3	Complex Formation between Chaperones and Polypeptide Substrates	183
5.6.4	Identification of Chaperone-binding Sites	184
5.6.5	Chaperone-mediated Refolding of Test Proteins	186
5.6.6	ATPase Activity	188
	Acknowledgments	188
	References	189
6	Physical Methods for Studies of Fiber Formation and Structure	197
	<i>Thomas Scheibel and Louise Serpell</i>	
6.1	Introduction	197
6.2	Overview: Protein Fibers Formed in Vivo	198
6.2.1	Amyloid Fibers	198
6.2.2	Silks	199
6.2.3	Collagens	199
6.2.4	Actin, Myosin, and Tropomyosin Filaments	200
6.2.5	Intermediate Filaments/Nuclear Lamina	202
6.2.6	Fibrinogen/Fibrin	203
6.2.7	Microtubules	203
6.2.8	Elastic Fibers	204
6.2.9	Flagella and Pili	204
6.2.10	Filamentary Structures in Rod-like Viruses	205
6.2.11	Protein Fibers Used by Viruses and Bacteriophages to Bind to Their Hosts	206
6.3	Overview: Fiber Structures	206

6.3.1	Study of the Structure of β -sheet-containing Proteins	207
6.3.1.1	Amyloid	207
6.3.1.2	Paired Helical Filaments	207
6.3.1.3	β -Silks	207
6.3.1.4	β -Sheet-containing Viral Fibers	208
6.3.2	α -Helix-containing Protein Fibers	209
6.3.2.1	Collagen	209
6.3.2.2	Tropomyosin	210
6.3.2.3	Intermediate Filaments	210
6.3.3	Protein Polymers Consisting of a Mixture of Secondary Structure	211
6.3.3.1	Tubulin	211
6.3.3.2	Actin and Myosin Filaments	212
6.4	Methods to Study Fiber Assembly	213
6.4.1	Circular Dichroism Measurements for Monitoring Structural Changes Upon Fiber Assembly	213
6.4.1.1	Theory of CD	213
6.4.1.2	Experimental Guide to Measure CD Spectra and Structural Transition Kinetics	214
6.4.2	Intrinsic Fluorescence Measurements to Analyze Structural Changes	215
6.4.2.1	Theory of Protein Fluorescence	215
6.4.2.2	Experimental Guide to Measure Trp Fluorescence	216
6.4.3	Covalent Fluorescent Labeling to Determine Structural Changes of Proteins with Environmentally Sensitive Fluorophores	217
6.4.3.1	Theory on Environmental Sensitivity of Fluorophores	217
6.4.3.2	Experimental Guide to Labeling Proteins With Fluorophores	218
6.4.4	1-Anilino-8-Naphthalensulfonate (ANS) Binding to Investigate Fiber Assembly	219
6.4.4.1	Theory on Using ANS Fluorescence for Detecting Conformational Changes in Proteins	219
6.4.4.2	Experimental Guide to Using ANS for Monitoring Protein Fiber Assembly	220
6.4.5	Light Scattering to Monitor Particle Growth	220
6.4.5.1	Theory of Classical Light Scattering	221
6.4.5.2	Theory of Dynamic Light Scattering	221
6.4.5.3	Experimental Guide to Analyzing Fiber Assembly Using DLS	222
6.4.6	Field-flow Fractionation to Monitor Particle Growth	222
6.4.6.1	Theory of FFF	222
6.4.6.2	Experimental Guide to Using FFF for Monitoring Fiber Assembly	223
6.4.7	Fiber Growth-rate Analysis Using Surface Plasmon Resonance	223
6.4.7.1	Theory of SPR	223
6.4.7.2	Experimental Guide to Using SPR for Fiber-growth Analysis	224
6.4.8	Single-fiber Growth Imaging Using Atomic Force Microscopy	225

6.4.8.1	Theory of Atomic Force Microscopy	225
6.4.8.2	Experimental Guide for Using AFM to Investigate Fiber Growth	225
6.4.9	Dyes Specific for Detecting Amyloid Fibers	226
6.4.9.1	Theory on Congo Red and Thioflavin T Binding to Amyloid	226
6.4.9.2	Experimental Guide to Detecting Amyloid Fibers with CR and Thioflavin Binding	227
6.5	Methods to Study Fiber Morphology and Structure	228
6.5.1	Scanning Electron Microscopy for Examining the Low-resolution Morphology of a Fiber Specimen	228
6.5.1.1	Theory of SEM	228
6.5.1.2	Experimental Guide to Examining Fibers by SEM	229
6.5.2	Transmission Electron Microscopy for Examining Fiber Morphology and Structure	230
6.5.2.1	Theory of TEM	230
6.5.2.2	Experimental Guide to Examining Fiber Samples by TEM	231
6.5.3	Cryo-electron Microscopy for Examination of the Structure of Fibrous Proteins	232
6.5.3.1	Theory of Cryo-electron Microscopy	232
6.5.3.2	Experimental Guide to Preparing Proteins for Cryo-electron Microscopy	233
6.5.3.3	Structural Analysis from Electron Micrographs	233
6.5.4	Atomic Force Microscopy for Examining the Structure and Morphology of Fibrous Proteins	234
6.5.4.1	Experimental Guide for Using AFM to Monitor Fiber Morphology	234
6.5.5	Use of X-ray Diffraction for Examining the Structure of Fibrous Proteins	236
6.5.5.1	Theory of X-Ray Fiber Diffraction	236
6.5.5.2	Experimental Guide to X-Ray Fiber Diffraction	237
6.5.6	Fourier Transformed Infrared Spectroscopy	239
6.5.6.1	Theory of FTIR	239
6.5.6.2	Experimental Guide to Determining Protein Conformation by FTIR	240
6.6	Concluding Remarks	241
	Acknowledgements	242
	References	242
7	Protein Unfolding in the Cell	254
	<i>Prakash Koodathingal, Neil E. Jaffe, and Andreas Matouschek</i>	
7.1	Introduction	254
7.2	Protein Translocation Across Membranes	254
7.2.1	Compartmentalization and Unfolding	254
7.2.2	Mitochondria Actively Unfold Precursor Proteins	256
7.2.3	The Protein Import Machinery of Mitochondria	257
7.2.4	Specificity of Unfolding	259

7.2.5	Protein Import into Other Cellular Compartments	259
7.3	Protein Unfolding and Degradation by ATP-dependent Proteases	260
7.3.1	Structural Considerations of Unfoldases Associated With Degradation	260
7.3.2	Unfolding Is Required for Degradation by ATP-dependent Proteases	261
7.3.3	The Role of ATP and Models of Protein Unfolding	262
7.3.4	Proteins Are Unfolded Sequentially and Processively	263
7.3.5	The Influence of Substrate Structure on the Degradation Process	264
7.3.6	Unfolding by Pulling	264
7.3.7	Specificity of Degradation	265
7.4	Conclusions	266
7.5	Experimental Protocols	266
7.5.1	Size of Import Channels in the Outer and Inner Membranes of Mitochondria	266
7.5.2	Structure of Precursor Proteins During Import into Mitochondria	266
7.5.3	Import of Barnase Mutants	267
7.5.4	Protein Degradation by ATP-dependent Proteases	267
7.5.5	Use of Multi-domain Substrates	268
7.5.6	Studies Using Circular Permutants	268
	References	269
8	Natively Disordered Proteins	275
	<i>Gary W. Daughdrill, Gary J. Pielak, Vladimir N. Uversky, Marc S. Cortese, and A. Keith Dunker</i>	
8.1	Introduction	275
8.1.1	The Protein Structure-Function Paradigm	275
8.1.2	Natively Disordered Proteins	277
8.1.3	A New Protein Structure-Function Paradigm	280
8.2	Methods Used to Characterize Natively Disordered Proteins	281
8.2.1	NMR Spectroscopy	281
8.2.1.1	Chemical Shifts Measure the Presence of Transient Secondary Structure	282
8.2.1.2	Pulsed Field Gradient Methods to Measure Translational Diffusion	284
8.2.1.3	NMR Relaxation and Protein Flexibility	284
8.2.1.4	Using the Model-free Analysis of Relaxation Data to Estimate Internal Mobility and Rotational Correlation Time	285
8.2.1.5	Using Reduced Spectral Density Mapping to Assess the Amplitude and Frequencies of Intramolecular Motion	286
8.2.1.6	Characterization of the Dynamic Structures of Natively Disordered Proteins Using NMR	287
8.2.2	X-ray Crystallography	288
8.2.3	Small Angle X-ray Diffraction and Hydrodynamic Measurements	293

8.2.4	Circular Dichroism Spectropolarimetry	297
8.2.5	Infrared and Raman Spectroscopy	299
8.2.6	Fluorescence Methods	301
8.2.6.1	Intrinsic Fluorescence of Proteins	301
8.2.6.2	Dynamic Quenching of Fluorescence	302
8.2.6.3	Fluorescence Polarization and Anisotropy	303
8.2.6.4	Fluorescence Resonance Energy Transfer	303
8.2.6.5	ANS Fluorescence	305
8.2.7	Conformational Stability	308
8.2.7.1	Effect of Temperature on Proteins with Extended Disorder	309
8.2.7.2	Effect of pH on Proteins with Extended Disorder	309
8.2.8	Mass Spectrometry-based High-resolution Hydrogen-Deuterium Exchange	309
8.2.9	Protease Sensitivity	311
8.2.10	Prediction from Sequence	313
8.2.11	Advantage of Multiple Methods	314
8.3	Do Natively Disordered Proteins Exist Inside Cells?	315
8.3.1	Evolution of Ordered and Disordered Proteins Is Fundamentally Different	315
8.3.1.1	The Evolution of Natively Disordered Proteins	315
8.3.1.2	Adaptive Evolution and Protein Flexibility	317
8.3.1.3	Phylogeny Reconstruction and Protein Structure	318
8.3.2	Direct Measurement by NMR	320
8.4	Functional Repertoire	322
8.4.1	Molecular Recognition	322
8.4.1.1	The Coupling of Folding and Binding	322
8.4.1.2	Structural Plasticity for the Purpose of Functional Plasticity	323
8.4.1.3	Systems Where Disorder Increases Upon Binding	323
8.4.2	Assembly/Disassembly	325
8.4.3	Highly Entropic Chains	325
8.4.4	Protein Modification	327
8.5	Importance of Disorder for Protein Folding	328
8.6	Experimental Protocols	331
8.6.1	NMR Spectroscopy	331
8.6.1.1	General Requirements	331
8.6.1.2	Measuring Transient Secondary Structure in Secondary Chemical Shifts	332
8.6.1.3	Measuring the Translational Diffusion Coefficient Using Pulsed Field Gradient Diffusion Experiments	332
8.6.1.4	Relaxation Experiments	332
8.6.1.5	Relaxation Data Analysis Using Reduced Spectral Density Mapping	333
8.6.1.6	In-cell NMR	334
8.6.2	X-ray Crystallography	334
8.6.3	Circular Dichroism Spectropolarimetry	336

Acknowledgements 337

References 337

9 The Catalysis of Disulfide Bond Formation in Prokaryotes 358

Jean-Francois Collet and James C. Bardwell

- 9.1 Introduction 358
- 9.2 Disulfide Bond Formation in the *E. coli* Periplasm 358
 - 9.2.1 A Small Bond, a Big Effect 358
 - 9.2.2 Disulfide Bond Formation Is a Catalyzed Process 359
 - 9.2.3 DsbA, a Protein-folding Catalyst 359
 - 9.2.4 How is DsbA Re-oxidized? 361
 - 9.2.5 From Where Does the Oxidative Power of DsbB Originate? 361
 - 9.2.6 How Are Disulfide Bonds Transferred From DsbB to DsbA? 362
 - 9.2.7 How Can DsbB Generate Disulfide by Quinone Reduction? 364
- 9.3 Disulfide Bond Isomerization 365
 - 9.3.1 The Protein Disulfide Isomerases DsbC and DsbG 365
 - 9.3.2 Dimerization of DsbC and DsbG Is Important for Isomerase and Chaperone Activity 366
 - 9.3.3 Dimerization Protects from DsbB Oxidation 367
 - 9.3.4 Import of Electrons from the Cytoplasm: DsbD 367
 - 9.3.5 Conclusions 369
- 9.4 Experimental Protocols 369
 - 9.4.1 Oxidation-reduction of a Protein Sample 369
 - 9.4.2 Determination of the Free Thiol Content of a Protein 370
 - 9.4.3 Separation by HPLC 371
 - 9.4.4 Tryptophan Fluorescence 372
 - 9.4.5 Assay of Disulfide Oxidase Activity 372
- References 373

10 Catalysis of Peptidyl-prolyl *cis/trans* Isomerization by Enzymes 377

Gunter Fischer

- 10.1 Introduction 377
- 10.2 Peptidyl-prolyl *cis/trans* Isomerization 379
- 10.3 Monitoring Peptidyl-prolyl *cis/trans* Isomerase Activity 383
- 10.4 Prototypical Peptidyl-prolyl *cis/trans* Isomerases 388
 - 10.4.1 General Considerations 388
 - 10.4.2 Prototypic Cyclophilins 390
 - 10.4.3 Prototypic FK506-binding Proteins 394
 - 10.4.4 Prototypic Parvulins 397
- 10.5 Concluding Remarks 399
- 10.6 Experimental Protocols 399
 - 10.6.1 PPIase Assays: Materials 399
 - 10.6.2 PPIase Assays: Equipment 400
 - 10.6.3 Assaying Procedure: Protease-coupled Spectrophotometric Assay 400

10.6.4	Assaying Procedure: Protease-free Spectrophotometric Assay	401
	References	401
11	Secondary Amide Peptide Bond <i>cis/trans</i> Isomerization in Polypeptide Backbone Restructuring: Implications for Catalysis	415
	<i>Cordelia Schiene-Fischer and Christian Lücke</i>	
11.1	Introduction	415
11.2	Monitoring Secondary Amide Peptide Bond <i>cis/trans</i> Isomerization	416
11.3	Kinetics and Thermodynamics of Secondary Amide Peptide Bond <i>cis/trans</i> Isomerization	418
11.4	Principles of DnaK Catalysis	420
11.5	Concluding Remarks	423
11.6	Experimental Protocols	424
11.6.1	Stopped-flow Measurements of Peptide Bond <i>cis/trans</i> Isomerization	424
11.6.2	Two-dimensional ¹ H-NMR Exchange Experiments	425
	References	426
12	Ribosome-associated Proteins Acting on Newly Synthesized Polypeptide Chains	429
	<i>Sabine Rospert, Matthias Gautschi, Magdalena Rakwalska, and Uta Raue</i>	
12.1	Introduction	429
12.2	Signal Recognition Particle, Nascent Polypeptide-associated Complex, and Trigger Factor	432
12.2.1	Signal Recognition Particle	432
12.2.2	An Interplay between Eukaryotic SRP and Nascent Polypeptide-associated Complex?	435
12.2.3	Interplay between Bacterial SRP and Trigger Factor?	435
12.2.4	Functional Redundancy: TF and the Bacterial Hsp70 Homologue DnaK	436
12.3	Chaperones Bound to the Eukaryotic Ribosome: Hsp70 and Hsp40 Systems	436
12.3.1	Sis1p and Ssa1p: an Hsp70/Hsp40 System Involved in Translation Initiation?	437
12.3.2	Ssb1/2p, an Hsp70 Homologue Distributed Between Ribosomes and Cytosol	438
12.3.3	Function of Ssb1/2p in Degradation and Protein Folding	439
12.3.4	Zuotin and Ssz1p: a Stable Chaperone Complex Bound to the Yeast Ribosome	440
12.3.5	A Functional Chaperone Triad Consisting of Ssb1/2p, Ssz1p, and Zuotin	440
12.3.6	Effects of Ribosome-bound Chaperones on the Yeast Prion [PSI ⁺]	442
12.4	Enzymes Acting on Nascent Polypeptide Chains	443

12.4.1	Methionine Aminopeptidases	443
12.4.2	<i>N</i> ² -acetyltransferases	444
12.5	A Complex Arrangement at the Yeast Ribosomal Tunnel Exit	445
12.6	Experimental Protocols	446
12.6.1	Purification of Ribosome-associated Protein Complexes from Yeast	446
12.6.2	Growth of Yeast and Preparation of Ribosome-associated Proteins by High-salt Treatment of Ribosomes	447
12.6.3	Purification of NAC and RAC	448
	References	449

Part II, Volume 2

13	The Role of Trigger Factor in Folding of Newly Synthesized Proteins	459
	<i>Elke Deuerling, Thomas Rauch, Holger Patzelt, and Bernd Bukau</i>	
13.1	Introduction	459
13.2	In Vivo Function of Trigger Factor	459
13.2.1	Discovery	459
13.2.2	Trigger Factor Cooperates With the DnaK Chaperone in the Folding of Newly Synthesized Cytosolic Proteins	460
13.2.3	In Vivo Substrates of Trigger Factor and DnaK	461
13.2.4	Substrate Specificity of Trigger Factor	463
13.3	Structure–Function Analysis of Trigger Factor	465
13.3.1	Domain Structure and Conservation	465
13.3.2	Quaternary Structure	468
13.3.3	PPIase and Chaperone Activity of Trigger Factor	469
13.3.4	Importance of Ribosome Association	470
13.4	Models of the Trigger Factor Mechanism	471
13.5	Experimental Protocols	473
13.5.1	Trigger Factor Purification	473
13.5.2	GAPDH Trigger Factor Activity Assay	475
13.5.3	Modular Cell-free <i>E. coli</i> Transcription/Translation System	475
13.5.4	Isolation of Ribosomes and Add-back Experiments	483
13.5.5	Cross-linking Techniques	485
	References	485
14	Cellular Functions of Hsp70 Chaperones	490
	<i>Elizabeth A. Craig and Peggy Huang</i>	
14.1	Introduction	490
14.2	“Soluble” Hsp70s/J-proteins Function in General Protein Folding	492
14.2.1	The Soluble Hsp70 of <i>E. coli</i> , DnaK	492
14.2.2	Soluble Hsp70s of Major Eukaryotic Cellular Compartments	493
14.2.2.1	Eukaryotic Cytosol	493
14.2.2.2	Matrix of Mitochondria	494
14.2.2.3	Lumen of the Endoplasmic Reticulum	494

14.3	“Tethered” Hsp70s/J-proteins: Roles in Protein Folding on the Ribosome and in Protein Translocation	495
14.3.1	Membrane-tethered Hsp70/J-protein	495
14.3.2	Ribosome-associated Hsp70/J-proteins	496
14.4	Modulating of Protein Conformation by Hsp70s/J-proteins	498
14.4.1	Assembly of Fe/S Centers	499
14.4.2	Uncoating of Clathrin-coated Vesicles	500
14.4.3	Regulation of the Heat Shock Response	501
14.4.4	Regulation of Activity of DNA Replication-initiator Proteins	502
14.5	Cases of a Single Hsp70 Functioning With Multiple J-Proteins	504
14.6	Hsp70s/J-proteins – When an Hsp70 Maybe Isn’t Really a Chaperone	504
14.6.1	The Ribosome-associated “Hsp70” Ssz1	505
14.6.2	Mitochondrial Hsp70 as the Regulatory Subunit of an Endonuclease	506
14.7	Emerging Concepts and Unanswered Questions	507
	References	507
15	Regulation of Hsp70 Chaperones by Co-chaperones	516
	<i>Matthias P. Mayer and Bernd Bukau</i>	
15.1	Introduction	516
15.2	Hsp70 Proteins	517
15.2.1	Structure and Conservation	517
15.2.2	ATPase Cycle	519
15.2.3	Structural Investigations	521
15.2.4	Interactions With Substrates	522
15.3	J-domain Protein Family	526
15.3.1	Structure and Conservation	526
15.3.2	Interaction With Hsp70s	530
15.3.3	Interactions with Substrates	532
15.4	Nucleotide Exchange Factors	534
15.4.1	GrpE: Structure and Interaction with DnaK	534
15.4.2	Nucleotide Exchange Reaction	535
15.4.3	Bag Family: Structure and Interaction With Hsp70	536
15.4.4	Relevance of Regulated Nucleotide Exchange for Hsp70s	538
15.5	TPR Motifs Containing Co-chaperones of Hsp70	540
15.5.1	Hip	541
15.5.2	Hop	542
15.5.3	Chip	543
15.6	Concluding Remarks	544
15.7	Experimental Protocols	544
15.7.1	Hsp70s	544
15.7.2	J-Domain Proteins	545
15.7.3	GrpE	546
15.7.4	Bag-1	547

15.7.5	Hip	548
15.7.6	Hop	549
15.7.7	Chip	549
	References	550
16	Protein Folding in the Endoplasmic Reticulum Via the Hsp70 Family	563
	<i>Ying Shen, Kyung Tae Chung, and Linda M. Hendershot</i>	
16.1	Introduction	563
16.2	BiP Interactions with Unfolded Proteins	564
16.3	ER-localized DnaJ Homologues	567
16.4	ER-localized Nucleotide-exchange/releasing Factors	571
16.5	Organization and Relative Levels of Chaperones in the ER	572
16.6	Regulation of ER Chaperone Levels	573
16.7	Disposal of BiP-associated Proteins That Fail to Fold or Assemble	575
16.8	Other Roles of BiP in the ER	576
16.9	Concluding Comments	576
16.10	Experimental Protocols	577
16.10.1	Production of Recombinant ER Proteins	577
16.10.1.1	General Concerns	577
16.10.1.2	Bacterial Expression	578
16.10.1.3	Yeast Expression	580
16.10.1.4	Baculovirus	581
16.10.1.5	Mammalian Cells	583
16.10.2	Yeast Two-hybrid Screen for Identifying Interacting Partners of ER Proteins	586
16.10.3	Methods for Determining Subcellular Localization, Topology, and Orientation of Proteins	588
16.10.3.1	Sequence Predictions	588
16.10.3.2	Immunofluorescence Staining	589
16.10.3.3	Subcellular Fractionation	589
16.10.3.4	Determination of Topology	590
16.10.3.5	N-linked Glycosylation	592
16.10.4	Nucleotide Binding, Hydrolysis, and Exchange Assays	594
16.10.4.1	Nucleotide-binding Assays	594
16.10.4.2	ATP Hydrolysis Assays	596
16.10.4.3	Nucleotide Exchange Assays	597
16.10.5	Assays for Protein-Protein Interactions in Vitro/in Vivo	599
16.10.5.1	In Vitro GST Pull-down Assay	599
16.10.5.2	Co-immunoprecipitation	600
16.10.5.3	Chemical Cross-linking	600
16.10.5.4	Yeast Two-hybrid System	601
16.10.6	In Vivo Folding, Assembly, and Chaperone-binding Assays	601
16.10.6.1	Monitoring Oxidation of Intrachain Disulfide Bonds	601
16.10.6.2	Detection of Chaperone Binding	602

Acknowledgements 603

References 603

17 Quality Control In Glycoprotein Folding 617

E. Sergio Trombetta and Armando J. Parodi

17.1 Introduction 617

17.2 ER N-glycan Processing Reactions 617

17.3 The UDP-Glc:Glycoprotein Glucosyltransferase 619

17.4 Protein Folding in the ER 621

17.5 Unconventional Chaperones (Lectins) Are Present in the ER Lumen 621

17.6 In Vivo Glycoprotein-CNX/CRT Interaction 623

17.7 Effect of CNX/CRT Binding on Glycoprotein Folding and ER Retention 624

17.8 Glycoprotein-CNX/CRT Interaction Is Not Essential for Unicellular Organisms and Cells in Culture 627

17.9 Diversion of Misfolded Glycoproteins to Proteasomal Degradation 629

17.10 Unfolding Irreparably Misfolded Glycoproteins to Facilitate Proteasomal Degradation 632

17.11 Summary and Future Directions 633

17.12 Characterization of N-glycans from Glycoproteins 634

17.12.1 Characterization of N-glycans Present in Immunoprecipitated Samples 634

17.12.2 Analysis of Radio-labeled N-glycans 636

17.12.3 Extraction and Analysis of Protein-bound N-glycans 636

17.12.4 GII and GT Assays 637

17.12.4.1 Assay for GII 637

17.12.4.2 Assay for GT 638

17.12.5 Purification of GII and GT from Rat Liver 639

References 641

18 Procollagen Biosynthesis in Mammalian Cells 649

Mohammed Tasab and Neil J. Bulleid

18.1 Introduction 649

18.1.1 Variety and Complexity of Collagen Proteins 649

18.1.2 Fibrillar Procollagen 650

18.1.3 Expression of Fibrillar Collagens 650

18.2 The Procollagen Biosynthetic Process: An Overview 651

18.3 Disulfide Bonding in Procollagen Assembly 653

18.4 The Influence of Primary Amino Acid Sequence on Intracellular Procollagen Folding 654

18.4.1 Chain Recognition and Type-specific Assembly 654

18.4.2 Assembly of Multi-subunit Proteins 654

18.4.3 Coordination of Type-specific Procollagen Assembly and Chain Selection 655

18.4.4	Hypervariable Motifs: Components of a Recognition Mechanism That Distinguishes Between Procollagen Chains?	656
18.4.5	Modeling the C-propeptide	657
18.4.6	Chain Association	657
18.5	Posttranslational Modifications That Affect Procollagen Folding	658
18.5.1	Hydroxylation and Triple-helix Stability	658
18.6	Procollagen Chaperones	658
18.6.1	Prolyl 4-Hydroxylase	658
18.6.2	Protein Disulfide Isomerase	659
18.6.3	Hsp47	660
18.6.4	PPI and BiP	661
18.7	Analysis of Procollagen Folding	662
18.8	Experimental Part	663
18.8.1	Materials Required	663
18.8.2	Experimental Protocols	664
	References	668
19	Redox Regulation of Chaperones	677
	<i>Jörg H. Hoffmann and Ursula Jakob</i>	
19.1	Introduction	677
19.2	Disulfide Bonds as Redox-Switches	677
19.2.1	Functionality of Disulfide Bonds	677
19.2.2	Regulatory Disulfide Bonds as Functional Switches	679
19.2.3	Redox Regulation of Chaperone Activity	680
19.3	Prokaryotic Hsp33: A Chaperone Activated by Oxidation	680
19.3.1	Identification of a Redox-regulated Chaperone	680
19.3.2	Activation Mechanism of Hsp33	681
19.3.3	The Crystal Structure of Active Hsp33	682
19.3.4	The Active Hsp33-Dimer: An Efficient Chaperone Holdase	683
19.3.5	Hsp33 is Part of a Sophisticated Multi-chaperone Network	684
19.4	Eukaryotic Protein Disulfide Isomerase (PDI): Redox Shuffling in the ER	685
19.4.1	PDI, A Multifunctional Enzyme in Eukaryotes	685
19.4.2	PDI and Redox Regulation	687
19.5	Concluding Remarks and Outlook	688
19.6	Appendix – Experimental Protocols	688
19.6.1	How to Work With Redox-regulated Chaperones in Vitro	689
19.6.1.1	Preparation of the Reduced Protein Species	689
19.6.1.2	Preparation of the Oxidized Protein Species	690
19.6.1.3	In Vitro Thiol Trapping to Monitor the Redox State of Proteins	691
19.6.2	Thiol Coordinating Zinc Centers as Redox Switches	691
19.6.2.1	PAR-PMPs Assay to Quantify Zinc	691
19.6.2.2	Determination of Zinc-binding Constants	692
19.6.3	Functional Analysis of Redox-regulated Chaperones in Vitro/in Vivo	693
19.6.3.1	Chaperone Activity Assays	693

19.6.3.2	Manipulating and Analyzing Redox Conditions in Vivo	694
	Acknowledgements	694
	References	694
20	The <i>E. coli</i> GroE Chaperone	699
	<i>Steven G. Burston and Stefan Walter</i>	
20.1	Introduction	699
20.2	The Structure of GroEL	699
20.3	The Structure of GroEL-ATP	700
20.4	The Structure of GroES and its Interaction with GroEL	701
20.5	The Interaction Between GroEL and Substrate Polypeptides	702
20.6	GroEL is a Complex Allosteric Macromolecule	703
20.7	The Reaction Cycle of the GroE Chaperone	705
20.8	The Effect of GroE on Protein-folding Pathways	708
20.9	Future Perspectives	710
20.10	Experimental Protocols	710
	Acknowledgments	719
	References	719
21	Structure and Function of the Cytosolic Chaperonin CCT	725
	<i>José M. Valpuesta, José L. Carrascosa, and Keith R. Willison</i>	
21.1	Introduction	725
21.2	Structure and Composition of CCT	726
21.3	Regulation of CCT Expression	729
21.4	Functional Cycle of CCT	730
21.5	Folding Mechanism of CCT	731
21.6	Substrates of CCT	735
21.7	Co-chaperones of CCT	739
21.8	Evolution of CCT	741
21.9	Concluding Remarks	743
21.10	Experimental Protocols	743
21.10.1	Purification	743
21.10.2	ATP Hydrolysis Measurements	744
21.10.3	CCT Substrate-binding and Folding Assays	744
21.10.4	Electron Microscopy and Image Processing	744
	References	747
22	Structure and Function of GimC/Prefoldin	756
	<i>Katja Siegers, Andreas Bracher, and Ulrich Hartl</i>	
22.1	Introduction	756
22.2	Evolutionary Distribution of GimC/Prefoldin	757
22.3	Structure of the Archaeal GimC/Prefoldin	757
22.4	Complexity of the Eukaryotic/Archaeal GimC/Prefoldin	759
22.5	Functional Cooperation of GimC/Prefoldin With the Eukaryotic Chaperonin TRiC/CCT	761

22.6	Experimental Protocols	764
22.6.1	Actin-folding Kinetics	764
22.6.2	Prevention of Aggregation (Light-scattering) Assay	765
22.6.3	Actin-binding Assay	765
	Acknowledgements	766
	References	766
23	Hsp90: From Dispensable Heat Shock Protein to Global Player	768
	<i>Klaus Richter, Birgit Meinschmidt, and Johannes Buchner</i>	
23.1	Introduction	768
23.2	The Hsp90 Family in Vivo	768
23.2.1	Evolutionary Relationships within the Hsp90 Gene Family	768
23.2.2	In Vivo Functions of Hsp90	769
23.2.3	Regulation of Hsp90 Expression and Posttranscriptional Activation	772
23.2.4	Chemical Inhibition of Hsp90	773
23.2.5	Identification of Natural Hsp90 Substrates	774
23.3	In Vitro Investigation of the Chaperone Hsp90	775
23.3.1	Hsp90: A Special Kind of ATPase	775
23.3.2	The ATPase Cycle of Hsp90	780
23.3.3	Interaction of Hsp90 with Model Substrate Proteins	781
23.3.4	Investigating Hsp90 Substrate Interactions Using Native Substrates	783
23.4	Partner Proteins: Does Complexity Lead to Specificity?	784
23.4.1	Hop, p23, and PPIases: The Chaperone Cycle of Hsp90	784
23.4.2	Hop/Sti1: Interactions Mediated by TPR Domains	787
23.4.3	p23/Sba1: Nucleotide-specific Interaction with Hsp90	789
23.4.4	Large PPIases: Conferring Specificity to Substrate Localization?	790
23.4.5	Pp5: Facilitating Dephosphorylation	791
23.4.6	Cdc37: Building Complexes with Kinases	792
23.4.7	Tom70: Chaperoning Mitochondrial Import	793
23.4.8	CHIP and Sgt1: Multiple Connections to Protein Degradation	793
23.4.9	Aha1 and Hch1: Just Stimulating the ATPase?	794
23.4.10	Cns1, Sgt2, and Xap2: Is a TPR Enough to Become an Hsp90 Partner?	796
23.5	Outlook	796
23.6	Appendix – Experimental Protocols	797
23.6.1	Calculation of Phylogenetic Trees Based on Protein Sequences	797
23.6.2	Investigating the in Vivo Effect of Hsp90 Mutations in <i>S. cerevisiae</i>	797
23.6.3	Well-characterized Hsp90 Mutants	798
23.6.4	Investigating Activation of Heterologously Expressed Src Kinase in <i>S. cerevisiae</i>	800
23.6.5	Investigation of Heterologously Expressed Glucocorticoid Receptor in <i>S. cerevisiae</i>	800

23.6.6	Investigation of Chaperone Activity	801
23.6.7	Analysis of the ATPase Activity of Hsp90	802
23.6.8	Detecting Specific Influences on Hsp90 ATPase Activity	803
23.6.9	Investigation of the Quaternary Structure by SEC-HPLC	804
23.6.10	Investigation of Binding Events Using Changes of the Intrinsic Fluorescence	806
23.6.11	Investigation of Binding Events Using Isothermal Titration Calorimetry	807
23.6.12	Investigation of Protein-Protein Interactions Using Cross-linking	807
23.6.13	Investigation of Protein-Protein Interactions Using Surface Plasmon Resonance Spectroscopy	808
	Acknowledgements	810
	References	810
24	Small Heat Shock Proteins: Dynamic Players in the Folding Game	830
	<i>Franz Narberhaus and Martin Haslbeck</i>	
24.1	Introduction	830
24.2	α -Crystallins and the Small Heat Shock Protein Family: Diverse Yet Similar	830
24.3	Cellular Functions of α -Hsps	831
24.3.1	Chaperone Activity in Vitro	831
24.3.2	Chaperone Function in Vivo	835
24.3.3	Other Functions	836
24.4	The Oligomeric Structure of α -Hsps	837
24.5	Dynamic Structures as Key to Chaperone Activity	839
24.6	Experimental Protocols	840
24.6.1	Purification of sHsps	840
24.6.2	Chaperone Assays	843
24.6.3	Monitoring Dynamics of sHsps	846
	Acknowledgements	847
	References	848
25	Alpha-crystallin: Its Involvement in Suppression of Protein Aggregation and Protein Folding	858
	<i>Joseph Horwitz</i>	
25.1	Introduction	858
25.2	Distribution of Alpha-crystallin in the Various Tissues	858
25.3	Structure	859
25.4	Phosphorylation and Other Posttranslation Modification	860
25.5	Binding of Target Proteins to Alpha-crystallin	861
25.6	The Function of Alpha-crystallin	863
25.7	Experimental Protocols	863
25.7.1	Preparation of Alpha-crystallin	863
	Acknowledgements	870
	References	870

26	Transmembrane Domains in Membrane Protein Folding, Oligomerization, and Function	876
	<i>Anja Ridder and Dieter Langosch</i>	
26.1	Introduction	876
26.1.1	Structure of Transmembrane Domains	876
26.1.2	The Biosynthetic Route towards Folded and Oligomeric Integral Membrane Proteins	877
26.1.3	Structure and Stability of TMSs	878
26.1.3.1	Amino Acid Composition of TMSs and Flanking Regions	878
26.1.3.2	Stability of Transmembrane Helices	879
26.2	The Nature of Transmembrane Helix-Helix Interactions	880
26.2.1	General Considerations	880
26.2.1.1	Attractive Forces within Lipid Bilayers	880
26.2.1.2	Forces between Transmembrane Helices	881
26.2.1.3	Entropic Factors Influencing Transmembrane Helix-Helix Interactions	882
26.2.2	Lessons from Sequence Analyses and High-resolution Structures	883
26.2.3	Lessons from Bitopic Membrane Proteins	886
26.2.3.1	Transmembrane Segments Forming Right-handed Pairs	886
26.2.3.2	Transmembrane Segments Forming Left-handed Assemblies	889
26.2.4	Selection of Self-interacting TMSs from Combinatorial Libraries	892
26.2.5	Role of Lipids in Packing/Assembly of Membrane Proteins	893
26.3	Conformational Flexibility of Transmembrane Segments	895
26.4	Experimental Protocols	897
26.4.1	Biochemical and Biophysical Techniques	897
26.4.1.1	Visualization of Oligomeric States by Electrophoretic Techniques	898
26.4.1.2	Hydrodynamic Methods	899
26.4.1.3	Fluorescence Resonance Transfer	900
26.4.2	Genetic Assays	901
26.4.2.1	The ToxR System	901
26.4.2.2	Other Genetic Assays	902
26.4.3	Identification of TMS-TMS Interfaces by Mutational Analysis	903
	References	904

Part II, Volume 3

27	SecB	919
	<i>Arnold J. M. Driessen, Janny de Wit, and Nico Nouwen</i>	
27.1	Introduction	919
27.2	Selective Binding of Preproteins by SecB	920
27.3	SecA-SecB Interaction	925
27.4	Preprotein Transfer from SecB to SecA	928
27.5	Concluding Remarks	929
27.6	Experimental Protocols	930
27.6.1	How to Analyze SecB-Preprotein Interactions	930

27.6.2	How to Analyze SecB-SecA Interaction	931
	Acknowledgements	932
	References	933
28	Protein Folding in the Periplasm and Outer Membrane of <i>E. coli</i>	938
	<i>Michael Ehrmann</i>	
28.1	Introduction	938
28.2	Individual Cellular Factors	940
28.2.1	The Proline Isomerases FkpA, PpiA, SurA, and PpiD	941
28.2.1.1	FkpA	942
28.2.1.2	PpiA	942
28.2.1.3	SurA	943
28.2.1.4	PpiD	943
28.2.2	Skp	944
28.2.3	Proteases and Protease/Chaperone Machines	945
28.2.3.1	The HtrA Family of Serine Proteases	946
28.2.3.2	<i>E. coli</i> HtrAs	946
28.2.3.3	DegP and DegQ	946
28.2.3.4	DegS	947
28.2.3.5	The Structure of HtrA	947
28.2.3.6	Other Proteases	948
28.3	Organization of Folding Factors into Pathways and Networks	950
28.3.1	Synthetic Lethality and Extragenic High-copy Suppressors	950
28.3.2	Reconstituted in Vitro Systems	951
28.4	Regulation	951
28.4.1	The Sigma E Pathway	951
28.4.2	The Cpx Pathway	952
28.4.3	The Bae Pathway	953
28.5	Future Perspectives	953
28.6	Experimental Protocols	954
28.6.1	Pulse Chase Immunoprecipitation	954
	Acknowledgements	957
	References	957
29	Formation of Adhesive Pili by the Chaperone-Usher Pathway	965
	<i>Michael Vetsch and Rudi Glockshuber</i>	
29.1	Basic Properties of Bacterial, Adhesive Surface Organelles	965
29.2	Structure and Function of Pilus Chaperones	970
29.3	Structure and Folding of Pilus Subunits	971
29.4	Structure and Function of Pilus Ushers	973
29.5	Conclusions and Outlook	976
29.6	Experimental Protocols	977
29.6.1	Test for the Presence of Type 1 Piliated <i>E. coli</i> Cells	977
29.6.2	Functional Expression of Pilus Subunits in the <i>E. coli</i> Periplasm	977
29.6.3	Purification of Pilus Subunits from the <i>E. coli</i> Periplasm	978

29.6.4	Preparation of Ushers	979
	Acknowledgements	979
	References	980

30 Unfolding of Proteins During Import into Mitochondria 987

Walter Neupert, Michael Brunner, and Kai Hell

30.1	Introduction	987
30.2	Translocation Machineries and Pathways of the Mitochondrial Protein Import System	988
30.2.1	Import of Proteins Destined for the Mitochondrial Matrix	990
30.3	Import into Mitochondria Requires Protein Unfolding	993
30.4	Mechanisms of Unfolding by the Mitochondrial Import Motor	995
30.4.1	Targeted Brownian Ratchet	995
30.4.2	Power-stroke Model	995
30.5	Studies to Discriminate between the Models	996
30.5.1	Studies on the Unfolding of Preproteins	996
30.5.1.1	Comparison of the Import of Folded and Unfolded Proteins	996
30.5.1.2	Import of Preproteins With Different Presequence Lengths	999
30.5.1.3	Import of Titin Domains	1000
30.5.1.4	Unfolding by the Mitochondrial Membrane Potential $\Delta\Psi$	1000
30.5.2	Mechanistic Studies of the Import Motor	1000
30.5.2.1	Brownian Movement of the Polypeptide Within the Import Channel	1000
30.5.2.2	Recruitment of mtHsp70 by Tim44	1001
30.5.2.3	Import Without Recruitment of mtHsp70 by Tim44	1002
30.5.2.4	MtHsp70 Function in the Import Motor	1003
30.6	Discussion and Perspectives	1004
30.7	Experimental Protocols	1006
30.7.1	Protein Import Into Mitochondria in Vitro	1006
30.7.2	Stabilization of the DHFR Domain by Methotrexate	1008
30.7.3	Import of Precursor Proteins Unfolded With Urea	1009
30.7.4	Kinetic Analysis of the Unfolding Reaction by Trapping of Intermediates	1009
	References	1011

31 The Chaperone System of Mitochondria 1020

Wolfgang Voos and Nikolaus Pfanner

31.1	Introduction	1020
31.2	Membrane Translocation and the Hsp70 Import Motor	1020
31.3	Folding of Newly Imported Proteins Catalyzed by the Hsp70 and Hsp60 Systems	1026
31.4	Mitochondrial Protein Synthesis and the Assembly Problem	1030
31.5	Aggregation versus Degradation: Chaperone Functions Under Stress Conditions	1033
31.6	Experimental Protocols	1034

31.6.1	Chaperone Functions Characterized With Yeast Mutants	1034
31.6.2	Interaction of Imported Proteins With Matrix Chaperones	1036
31.6.3	Folding of Imported Model Proteins	1037
31.6.4	Assaying Mitochondrial Degradation of Imported Proteins	1038
31.6.5	Aggregation of Proteins in the Mitochondrial Matrix	1038
	References	1039
32	Chaperone Systems in Chloroplasts	1047
	<i>Thomas Becker, Jürgen Soll, and Enrico Schleiff</i>	
32.1	Introduction	1047
32.2	Chaperone Systems within Chloroplasts	1048
32.2.1	The Hsp70 System of Chloroplasts	1048
32.2.1.1	The Chloroplast Hsp70s	1049
32.2.1.2	The Co-chaperones of Chloroplastic Hsp70s	1051
32.2.2	The Chaperonins	1052
32.2.3	The HSP100/Clp Protein Family in Chloroplasts	1056
32.2.4	The Small Heat Shock Proteins	1058
32.2.5	Hsp90 Proteins of Chloroplasts	1061
32.2.6	Chaperone-like Proteins	1062
32.2.6.1	The Protein Disulfide Isomerase (PDI)	1062
32.2.6.2	The Peptidyl-prolyl <i>cis</i> Isomerase (PPIase)	1063
32.3	The Functional Chaperone Pathways in Chloroplasts	1065
32.3.1	Chaperones Involved in Protein Translocation	1065
32.3.2	Protein Transport Inside of Plastids	1070
32.3.3	Protein Folding and Complex Assembly Within Chloroplasts	1071
32.3.4	Chloroplast Chaperones Involved in Proteolysis	1072
32.3.5	Protein Storage Within Plastids	1073
32.3.6	Protein Protection and Repair	1074
32.4	Experimental Protocols	1075
32.4.1	Characterization of Cpn60 Binding to the Large Subunit of Rubisco via Native PAGE (adopted from Ref. [6])	1075
32.4.2	Purification of Chloroplast Cpn60 From Young Pea Plants (adopted from Ref. [203])	1076
32.4.3	Purification of Chloroplast Hsp21 From Pea (<i>Pisum sativum</i>) (adopted from [90])	1077
32.4.4	Light-scattering Assays for Determination of the Chaperone Activity Using Citrate Synthase as Substrate (adopted from [196])	1078
32.4.5	The Use Of Bis-ANS to Assess Surface Exposure of Hydrophobic Domains of Hsp17 of <i>Synechocystis</i> (adopted from [202])	1079
32.4.6	Determination of Hsp17 Binding to Lipids (adopted from Refs. [204, 205])	1079
	References	1081
33	An Overview of Protein Misfolding Diseases	1093
	<i>Christopher M. Dobson</i>	
33.1	Introduction	1093

33.2	Protein Misfolding and Its Consequences for Disease	1094
33.3	The Structure and Mechanism of Amyloid Formation	1097
33.4	A Generic Description of Amyloid Formation	1101
33.5	The Fundamental Origins of Amyloid Disease	1104
33.6	Approaches to Therapeutic Intervention in Amyloid Disease	1106
33.7	Concluding Remarks	1108
	Acknowledgements	1108
	References	1109
34	Biochemistry and Structural Biology of Mammalian Prion Disease	1114
	<i>Rudi Glockshuber</i>	
34.1	Introduction	1114
34.1.1	Prions and the “Protein-Only” Hypothesis	1114
34.1.2	Models of PrP ^{Sc} Propagation	1115
34.2	Properties of PrP ^C and PrP ^{Sc}	1117
34.3	Three-dimensional Structure and Folding of Recombinant PrP	1120
34.3.1	Expression of the Recombinant Prion Protein for Structural and Biophysical Studies	1120
34.3.2	Three-dimensional Structures of Recombinant Prion Proteins from Different Species and Their Implications for the Species Barrier of Prion Transmission	1120
34.3.2.1	Solution Structure of Murine PrP	1120
34.3.2.2	Comparison of Mammalian Prion Protein Structures and the Species Barrier of Prion Transmission	1124
34.3.3	Biophysical Characterization of the Recombinant Prion Protein	1125
34.3.3.1	Folding and Stability of Recombinant PrP	1125
34.3.3.2	Role of the Disulfide Bond in PrP	1127
34.3.3.3	Influence of Point Mutations Linked With Inherited TSEs on the Stability of Recombinant PrP	1129
34.4	Generation of Infectious Prions in Vitro: Principal Difficulties in Proving the Protein-Only Hypothesis	1131
34.5	Understanding the Strain Phenomenon in the Context of the Protein-Only Hypothesis: Are Prions Crystals?	1132
34.6	Conclusions and Outlook	1135
34.7	Experimental Protocols	1136
34.7.1	Protocol 1 [53, 55]	1136
34.7.2	Protocol 2 [54]	1137
	References	1138
35	Insights into the Nature of Yeast Prions	1144
	<i>Lev Z. Osherovich and Jonathan S. Weissman</i>	
35.1	Introduction	1144
35.2	Prions as Heritable Amyloidoses	1145
35.3	Prion Strains and Species Barriers: Universal Features of Amyloid-based Prion Elements	1149

35.4	Prediction and Identification of Novel Prion Elements	1151
35.5	Requirements for Prion Inheritance beyond Amyloid-mediated Growth	1154
35.6	Chaperones and Prion Replication	1157
35.7	The Structure of Prion Particles	1158
35.8	Prion-like Structures as Protein Interaction Modules	1159
35.9	Experimental Protocols	1160
35.9.1	Generation of Sup35 Amyloid Fibers in Vitro	1160
35.9.2	Thioflavin T-based Amyloid Seeding Efficacy Assay (Adapted from Chien et al. 2003)	1161
35.9.3	AFM-based Single-fiber Growth Assay	1162
35.9.4	Prion Infection Protocol (Adapted from Tanaka et al. 2004)	1164
35.9.5	Preparation of Lyticase	1165
35.9.6	Protocol for Counting Heritable Prion Units (Adapted from Cox et al. 2003)	1166
	Acknowledgements	1167
	References	1168
36	Polyglutamine Aggregates as a Model for Protein-misfolding Diseases	1175
	<i>Soojin Kim, James F. Morley, Anat Ben-Zvi, and Richard I. Morimoto</i>	
36.1	Introduction	1175
36.2	Polyglutamine Diseases	1175
36.2.1	Genetics	1175
36.2.2	Polyglutamine Diseases Involve a Toxic Gain of Function	1176
36.3	Polyglutamine Aggregates	1176
36.3.1	Presence of the Expanded Polyglutamine Is Sufficient to Induce Aggregation in Vivo	1176
36.3.2	Length of the Polyglutamine Dictates the Rate of Aggregate Formation	1177
36.3.3	Polyglutamine Aggregates Exhibit Features Characteristic of Amyloids	1179
36.3.4	Characterization of Protein Aggregates in Vivo Using Dynamic Imaging Methods	1180
36.4	A Role for Oligomeric Intermediates in Toxicity	1181
36.5	Consequences of Misfolded Proteins and Aggregates on Protein Homeostasis	1181
36.6	Modulators of Polyglutamine Aggregation and Toxicity	1184
36.6.1	Protein Context	1184
36.6.2	Molecular Chaperones	1185
36.6.3	Proteasomes	1188
36.6.4	The Protein-folding “Buffer” and Aging	1188
36.6.5	Summary	1189
36.7	Experimental Protocols	1190
36.7.1	FRAP Analysis	1190
	References	1192

37	Protein Folding and Aggregation in the Expanded Polyglutamine Repeat Diseases	1200
	<i>Ronald Wetzel</i>	
37.1	Introduction	1200
37.2	Key Features of the Polyglutamine Diseases	1201
37.2.1	The Variety of Expanded PolyGln Diseases	1201
37.2.2	Clinical Features	1201
37.2.2.1	Repeat Expansions and Repeat Length	1202
37.2.3	The Role of PolyGln and PolyGln Aggregates	1203
37.3	PolyGln Peptides in Studies of the Molecular Basis of Expanded Polyglutamine Diseases	1205
37.3.1	Conformational Studies	1205
37.3.2	Preliminary in Vitro Aggregation Studies	1206
37.3.3	In Vivo Aggregation Studies	1206
37.4	Analyzing Polyglutamine Behavior With Synthetic Peptides: Practical Aspects	1207
37.4.1	Disaggregation of Synthetic Polyglutamine Peptides	1209
37.4.2	Growing and Manipulating Aggregates	1210
37.4.2.1	Polyglutamine Aggregation by Freeze Concentration	1210
37.4.2.2	Preparing Small Aggregates	1211
37.5	In vitro Studies of PolyGln Aggregation	1212
37.5.1	The Universe of Protein Aggregation Mechanisms	1212
37.5.2	Basic Studies on Spontaneous Aggregation	1213
37.5.3	Nucleation Kinetics of PolyGln	1215
37.5.4	Elongation Kinetics	1218
37.5.4.1	Microtiter Plate Assay for Elongation Kinetics	1219
37.5.4.2	Repeat-length and Aggregate-size Dependence of Elongation Rates	1220
37.6	The Structure of PolyGln Aggregates	1221
37.6.1	Electron Microscopy Analysis	1222
37.6.2	Analysis with Amyloid Dyes Thioflavin T and Congo Red	1222
37.6.3	Circular Dichroism Analysis	1224
37.6.4	Presence of a Generic Amyloid Epitope in PolyGln Aggregates	1225
37.6.5	Proline Mutagenesis to Dissect the Polyglutamine Fold Within the Aggregate	1225
37.7	Polyglutamine Aggregates and Cytotoxicity	1227
37.7.1	Direct Cytotoxicity of PolyGln Aggregates	1228
37.7.1.1	Delivery of Aggregates into Cells and Cellular Compartments	1229
37.7.1.2	Cell Killing by Nuclear-targeted PolyGln Aggregates	1229
37.7.2	Visualization of Functional, Recruitment-positive Aggregation Foci	1230
37.8	Inhibitors of polyGln Aggregation	1231
37.8.1	Designed Peptide Inhibitors	1231
37.8.2	Screening for Inhibitors of PolyGln Elongation	1231
37.9	Concluding Remarks	1232
37.10	Experimental Protocols	1233

37.10.1	Disaggregation of Synthetic PolyGln Peptides	1233
37.10.2	Determining the Concentration of Low-molecular-weight PolyGln Peptides by HPLC	1235
	Acknowledgements	1237
	References	1238
38	Production of Recombinant Proteins for Therapy, Diagnostics, and Industrial Research by in Vitro Folding	1245
	<i>Christian Lange and Rainer Rudolph</i>	
38.1	Introduction	1245
38.1.1	The Inclusion Body Problem	1245
38.1.2	Cost and Scale Limitations in Industrial Protein Folding	1248
38.2	Treatment of Inclusion Bodies	1250
38.2.1	Isolation of Inclusion Bodies	1250
38.2.2	Solubilization of Inclusion Bodies	1250
38.3	Refolding in Solution	1252
38.3.1	Protein Design Considerations	1252
38.3.2	Oxidative Refolding With Disulfide Bond Formation	1253
38.3.3	Transfer of the Unfolded Proteins Into Refolding Buffer	1255
38.3.4	Refolding Additives	1257
38.3.5	Cofactors in Protein Folding	1260
38.3.6	Chaperones and Folding-helper Proteins	1261
38.3.7	An Artificial Chaperone System	1261
38.3.8	Pressure-induced Folding	1262
38.3.9	Temperature-leap Techniques	1263
38.3.10	Recycling of Aggregates	1264
38.4	Alternative Refolding Techniques	1264
38.4.1	Matrix-assisted Refolding	1264
38.4.2	Folding by Gel Filtration	1266
38.4.3	Direct Refolding of Inclusion Body Material	1267
38.5	Conclusions	1268
38.6	Experimental Protocols	1268
38.6.1	Protocol 1: Isolation of Inclusion Bodies	1268
38.6.2	Protocol 2: Solubilization of Inclusion Bodies	1269
38.6.3	Protocol 3: Refolding of Proteins	1270
	Acknowledgements	1271
	References	1271
39	Engineering Proteins for Stability and Efficient Folding	1281
	<i>Bernhard Schimmele and Andreas Plückthun</i>	
39.1	Introduction	1281
39.2	Kinetic and Thermodynamic Aspects of Natural Proteins	1281
39.2.1	The Stability of Natural Proteins	1281
39.2.2	Different Kinds of "Stability"	1282
39.2.2.1	Thermodynamic Stability	1283

39.2.2.2	Kinetic Stability	1285
39.2.2.3	Folding Efficiency	1287
39.3	The Engineering Approach	1288
39.3.1	Consensus Strategies	1288
39.3.1.1	Principles	1288
39.3.1.2	Examples	1291
39.3.2	Structure-based Engineering	1292
39.3.2.1	Entropic Stabilization	1294
39.3.2.2	Hydrophobic Core Packing	1296
39.3.2.3	Charge Interactions	1297
39.3.2.4	Hydrogen Bonding	1298
39.3.2.5	Disallowed Phi-Psi Angles	1298
39.3.2.6	Local Secondary Structure Propensities	1299
39.3.2.7	Exposed Hydrophobic Side Chains	1299
39.3.2.8	Inter-domain Interactions	1300
39.3.3	Case Study: Combining Consensus Design and Rational Engineering to Yield Antibodies with Favorable Biophysical Properties	1300
39.4	The Selection and Evolution Approach	1305
39.4.1	Principles	1305
39.4.2	Screening and Selection Technologies Available for Improving Biophysical Properties	1311
39.4.2.1	In Vitro Display Technologies	1313
39.4.2.2	Partial in Vitro Display Technologies	1314
39.4.2.3	In Vivo Selection Technologies	1315
39.4.3	Selection for Enhanced Biophysical Properties	1316
39.4.3.1	Selection for Solubility	1316
39.4.3.2	Selection for Protein Display Rates	1317
39.4.3.3	Selection on the Basis of Cellular Quality Control	1318
39.4.4	Selection for Increased Stability	1319
39.4.4.1	General Strategies	1319
39.4.4.2	Protein Destabilization	1319
39.4.4.3	Selections Based on Elevated Temperature	1321
39.4.4.4	Selections Based on Destabilizing Agents	1322
39.4.4.5	Selection for Proteolytic Stability	1323
39.5	Conclusions and Perspectives	1324
	Acknowledgements	1326
	References	1326